Thematic Tutorials *Release 6.10*

The Sage Development Team

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Here you will find Sage demonstrations, quick reference cards, primers, and thematic tutorials,

- A *quickref* (or quick reference card) is a one page document with the essential examples, and pointers to the main entry points.
- A *primer* is a document meant for a user to get started by himself on a theme in a matter of minutes.
- A tutorial is more in-depth and could take as much as an hour or more to get through.

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12.1.1 Algebraic Combinatorics in Sage

Author: Anne Schilling (UC Davis)

These notes provide some Sage examples for Stanley's book:

A free pdf version of the book without exercises can be found on Stanley's homepage.

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I would like to thank Federico Castillo (who wrote a first version of the *n*-cube section) and Nicolas M. Thiery (who wrote a slightly different French version of the Tsetlin library section) for their help.

Walks in graphs

This section provides some examples on Chapter 1 of Stanley's book [Stanley2013].

We begin by creating a graph with 4 vertices:

```
sage: G = Graph(4)
sage: G
Graph on 4 vertices
```

This graph has no edges yet:

```
sage: G.vertices()
[0, 1, 2, 3]
sage: G.edges()
[]
```

Before we can add edges, we need to tell Sage that our graph can have loops and multiple edges.:

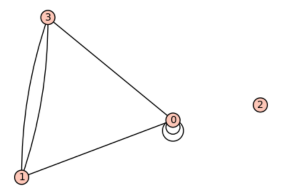
```
sage: G.allow_loops(True)
sage: G.allow_multiple_edges(True)
```

Now we are ready to add our edges by specifying a tuple of vertices that are connected by an edge. If there are multiple edges, we need to add the tuple with multiplicity.:

```
sage: G.add_edges([(0,0),(0,0),(0,1),(0,3),(1,3),(1,3)])
```

Now let us look at the graph!

```
sage: G.plot()
Graphics object consisting of 11 graphics primitives
```



We can construct the adjacency matrix:

```
sage: A = G.adjacency_matrix()
sage: A
[2 1 0 1]
[1 0 0 2]
[0 0 0 0]
[1 2 0 0]
```

The entry in row i and column j of the ℓ -th power of A gives us the number of paths of length ℓ from vertex i to vertex j. Let us verify this:

```
sage: A**2
[6 4 0 4]
[4 5 0 1]
[0 0 0 0]
[4 1 0 5]
```

There are 4 paths of length 2 from vertex 0 to vertex 1: take either loop at 0 and then the edge (0,1) (2 choices) or take the edge (0,3) and then either of the two edges (3,1) (two choices):

```
sage: (A**2)[0,1]
4
```

To count the number of closed walks, we can also look at the sum of the ℓ -th powers of the eigenvalues. Even though the eigenvalues are not integers, we find that the sum of their squares is an integer:

```
sage: A.eigenvalues()
[0, -2, 0.5857864376269049?, 3.414213562373095?]
sage: sum(la**2 for la in A.eigenvalues())
16.000000000000000?
```

We can achieve the same by looking at the trace of the ℓ -th power of the matrix:

```
sage: (A**2).trace()
16
```

n-Cube

This section provides some examples on Chapter 2 of Stanley's book [Stanley2013], which deals with n-cubes, the Radon transform, and combinatorial formulas for walks on the n-cube.

The vertices of the *n*-cube can be described by vectors in \mathbb{Z}_2^n . First we define the addition of two vectors $u, v \in \mathbb{Z}_2^n$ via the following distance:

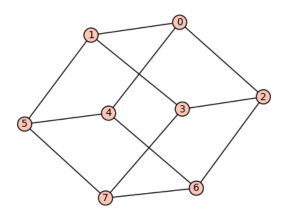
The distance function measures in how many slots two vectors in \mathbb{Z}_2^n differ:

```
sage: u=(1,0,1,1,1,0)
sage: v=(0,0,1,1,0,0)
sage: dist(u,v)
```

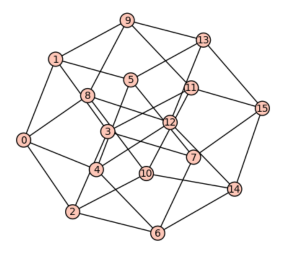
Now we are going to define the n-cube as the graph with vertices in \mathbb{Z}_2^n and edges between vertex u and vertex v if they differ in one slot, that is, the distance function is 1:

We can plot the 3 and 4-cube:

```
sage: cube(3).plot()
Graphics object consisting of 21 graphics primitives
```



```
sage: cube(4).plot()
Graphics object consisting of 49 graphics primitives
```



Next we can experiment and check Corollary 2.4 in Stanley's book, which states the n-cube has n choose i eigenvalues equal to n-2i:

```
sage: G = cube(2)
sage: G.adjacency_matrix().eigenvalues()
[2, -2, 0, 0]

sage: G = cube(3)
sage: G.adjacency_matrix().eigenvalues()
[3, -3, 1, 1, 1, -1, -1, -1]

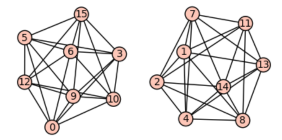
sage: G = cube(4)
sage: G.adjacency_matrix().eigenvalues()
[4, -4, 2, 2, 2, 2, -2, -2, -2, 0, 0, 0, 0, 0, 0]
```

It is now easy to slightly vary this problem and change the edge set by connecting vertices u and v if their distance is 2 (see Problem 4 in Chapter 2):

```
sage: def cube_2(n):
      G = Graph(2**n)
. . . . :
          vertices = Tuples([0,1],n)
. . . . :
         for i in range (2**n):
. . . . :
              for j in range (2**n):
                  if dist(vertices[i], vertices[j]) == 2:
                       G.add_edge(i, j)
. . . . :
         return G
sage: G = cube_2(2)
sage: G.adjacency_matrix().eigenvalues()
[1, 1, -1, -1]
sage: G = cube_2(4)
sage: G.adjacency_matrix().eigenvalues()
[6, 6, -2, -2, -2, -2, -2, -2, 0, 0, 0, 0, 0, 0, 0, 0]
```

Note that the graph is in fact disconnected. Do you understand why?

```
sage: cube_2(4).plot()
Graphics object consisting of 65 graphics primitives
```



The Tsetlin library

Introduction

In this section, we study a simple random walk (or Markov chain), called the *Tsetlin library*. It will give us the opportunity to see the interplay between combinatorics, linear algebra, representation theory and computer exploration, without requiring heavy theoretical background. I hope this encourages everyone to play around with this or similar systems and investigate their properties! Formal theorems and proofs can be found in the references at the end of this section.

It has been known for several years that the theory of group representations can facilitate the study of systems whose evolution is random (Markov chains), breaking them down into simpler systems. More recently it was realized that generalizing this (namely replacing the invertibility axiom for groups by other axioms) explains the behavior of other particularly simple Markov chains such as the Tsetlin library.

The Tsetlin library

Consider a bookshelf in a library containing n distinct books. When a person borrows a book and then returns it, it gets placed back on the shelf to the right of all books. This is what we naturally do with our pile of shirts in the closet: after use and cleaning, the shirt is placed on the top of its pile. Hence the most popular books/shirts will more likely appear on the right/top of the shelf/pile.

This type of organization has the advantage of being self-adaptive:

- The books most often used accumulate on the right and thus can easily be found.
- If the use changes over time, the system adapts.

In fact, this type of strategy is used not only in everyday life, but also in computer science. The natural questions that arise are:

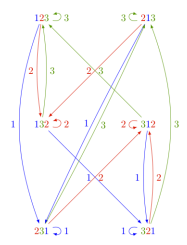
- Stationary distribution: To which state(s) does the system converge to? This, among other things, is used to evaluate the average access time to a book.
- The rate of convergence: How fast does the system adapt to a changing environment.

Let us formalize the description. The Tsetlin library is a discrete Markov chain (discrete time, discrete state space) described by:

- The state space Ω_n is given by the set of all permutations of the n books.
- The transition operators are denoted by $\partial_i \colon \Omega_n \to \Omega_n$. When ∂_i is applied to a permutation σ , the number i is moved to the end of the permutation.
- We assign parameters $x_i \ge 0$ for all $1 \le i \le n$ with $\sum_{i=1}^n x_i = 1$. The parameter x_i indicates the probability of choosing the operator ∂_i .

Transition graph and matrix

One can depict the action of the operators ∂_i on the state space Ω_n by a digraph. The following picture shows the action of $\partial_1, \partial_2, \partial_3$ on Ω_3 :



The above picture can be reproduced in Sage as follows:

```
sage: P = Poset(([1,2,3],[]))
```

This is the antichain poset. Its linear extensions are all permutations of $\{1, 2, 3\}$:

```
sage: L = P.linear_extensions()
sage: L
The set of all linear extensions of Finite poset containing 3 elements
sage: L.list()
[[3, 2, 1], [3, 1, 2], [2, 3, 1], [2, 1, 3], [1, 3, 2], [1, 2, 3]]
```

The graph is produced via:

```
sage: G = L.markov_chain_digraph(labeling='source'); G
Looped multi-digraph on 6 vertices
sage: view(G) # not tested
```

We can now look at the transition matrix and see whether we notice anything about its eigenvalue and eigenvectors:

```
sage: M = L.markov_chain_transition_matrix(labeling='source')
sage: M
                                              0]
             x0
[-x1 - x2]
                     0
                              0
                                     x0
      x1 - x0 - x2
                             0
                     x1
                                     0
                                              0 ]
      0 \quad 0 - x1 - x2
                            x0
                                     0
                                             x0]
      x2
             0
                     x2 - x0 - x1
                                     0
[
                     0 	 x1 - x0 - x2
Γ
[
      0
             x2
                     0
                              0 	 x2 - x0 - x1
```

This matrix is normalized so that all columns add to 0. So we need to add $(x_0 + x_1 + x_2)$ times the 6×6 identity matrix to get the probability matrix:

```
sage: x = M.base_ring().gens()
sage: Mt = (x[0]+x[1]+x[2])*matrix.identity(6)+M
sage: Mt
[x0 x0 0 0 x0 0]
[x1 x1 x1 0 0 0]
```

```
[ 0 0 x0 x0 0 x0]
[x2 0 x2 x2 0 0]
[ 0 0 0 x1 x1 x1]
[ 0 x2 0 0 x2 x2]
```

Since the x_i are formal variables, we need to compute the eigenvalues and eigenvectors in the symbolic ring SR:

```
sage: Mt.change_ring(SR).eigenvalues()
[x2, x1, x0, x0 + x1 + x2, 0, 0]
```

Do you see any pattern? In fact, if you start playing with bigger values of n (the size of the underlying permutations), you might observe that there is an eigenvalue for every subset S of $\{1, 2, \ldots, n\}$ and the multiplicity is given by a derangement number $d_{n-|S|}$. Derangment numbers count permutations without fixed point. For the eigenvectors we obtain:

```
sage: Mt.change_ring(SR).eigenvectors_right()
[(x2, [(0, 0, 0, 1, 0, -1)], 1),
   (x1, [(0, 1, 0, 0, -1, 0)], 1),
   (x0, [(1, 0, -1, 0, 0, 0)], 1),
   (x0 + x1 + x2,
   [(1, (x1 + x2)/(x0 + x2), x2/x1, (x1*x2 + x2^2)/(x0*x1 + x1^2),
        (x1*x2 + x2^2)/(x0^2 + x0*x2), (x1*x2 + x2^2)/(x0^2 + x0*x1))], 1),
   (0, [(1, 0, -1, 0, -1, 1), (0, 1, -1, 1, -1, 0)], 2)]
```

The stationary distribution is the eigenvector of eigenvalues $1 = x_0 + x_1 + x_2$. Do you see a pattern?

Optional exercices: Study of the transition operators and graph

Instead of using the methods that are already in Sage, try to build the state space Ω_n and the transition operators ∂_i yourself as follows.

1. For technical reasons, it is most practical in Sage to label the n books in the library by $0, 1, \dots, n-1$, and to represent each state in the Markov chain by a permutation of the set $\{0, \dots, n-1\}$ as a tuple. Construct the state space Ω_n as:

```
sage: map(tuple, Permutations(range(3)))
[(0, 1, 2), (0, 2, 1), (1, 0, 2), (1, 2, 0), (2, 0, 1), (2, 1, 0)]
```

- 2. Write a function transition_operator(sigma, i) which implements the operator ∂_i which takes as input a tuple sigma and integer $i \in \{1, 2, ..., n\}$ and outputs a new tuple. It might be useful to extract subtuples (sigma[i:j]) and concatentation.
- 3. Write a function tsetlin_digraph(n) which constructs the (multi digraph) as described as shown above. This can be achieved using DiGraph.
- 4. Verify for which values of n the digraph is strongly connected (i.e., you can go from any vertex to any other vertex by going in the direction of the arrow). This indicates whether the Markov chain is irreducible.

Conclusion

The Tsetlin library was studied from the viewpoint of monoids in [Bidigare1997] and [Brown2000]. Precise statements of the eigenvalues and the stationary distribution of the probability matrix as well as proofs of the statements are given in these papers. Generalizations of the Tsetlin library from the antichain to arbitrary posets was given in [AKS2013].

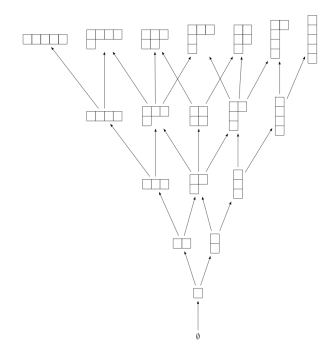
Young's lattice and the RSK algorithm

This section provides some examples on Young's lattice and the RSK (Robinson-Schensted-Knuth) algorithm explained in Chapter 8 of Stanley's book [Stanley2013].

Young's Lattice

We begin by creating the first few levels of Young's lattice Y. For this, we need to define the elements and the order relation for the poset, which is containment of partitions:

```
sage: level = 6
sage: elements = [b for n in range(level) for b in Partitions(n)]
sage: ord = lambda x,y: y.contains(x)
sage: Y = Poset((elements,ord), facade=True)
sage: H = Y.hasse_diagram()
sage: view(H) # optional - dot2tex graphviz
```



We can now define the up and down operators U and D on $\mathbf{Q}Y$. First we do so on partitions, which form a basis for $\mathbf{Q}Y$:

As a shorthand, one also can write the above as:

```
sage: U_on_basis = QQY.sum_of_monomials * Y.upper_covers
sage: D_on_basis = QQY.sum_of_monomials * Y.lower_covers
```

Here is the result when we apply the operators to the partition (2, 1):

```
sage: la = Partition([2,1])
sage: U_on_basis(la)
B[[2, 1, 1]] + B[[2, 2]] + B[[3, 1]]
sage: D_on_basis(la)
B[[1, 1]] + B[[2]]
```

Now we define the up and down operator on $\mathbf{Q}Y$:

```
sage: U = QQY.module_morphism(U_on_basis)
sage: D = QQY.module_morphism(D_on_basis)
```

We can check the identity $D_{i+1}U_i - U_{i-1}D_i = I_i$ explicitly on all partitions of i = 3:

```
sage: for p in Partitions(3):
....: b = QQY(p)
....: assert D(U(b)) - U(D(b)) == b
```

We can also check that the coefficient of $\lambda \vdash n$ in $U^n(\emptyset)$ is equal to the number of standard Young tableaux of shape λ :

```
sage: u = QQY(Partition([]))
sage: for i in range(4):
....: u = U(u)
sage: u
B[[1, 1, 1, 1]] + 3*B[[2, 1, 1]] + 2*B[[2, 2]] + 3*B[[3, 1]] + B[[4]]
```

For example, the number of standard Young tableaux of shape (2, 1, 1) is 3:

```
sage: StandardTableaux([2,1,1]).cardinality()
```

We can test this in general:

```
sage: for la in u.support():
....: assert u[la] == StandardTableaux(la).cardinality()
```

We can also check this against the hook length formula (Theorem 8.1):

RSK Algorithm

Let us now turn to the RSK algorithm. We can verify Example 8.12 as follows:

```
sage: p = Permutation([4,2,7,3,6,1,5])
sage: RSK(p)
[[[1, 3, 5], [2, 6], [4, 7]], [[1, 3, 5], [2, 4], [6, 7]]]
```

The tableaux can also be displayed as tableaux:

```
sage: P,Q = RSK(p)
sage: P.pp()
1  3  5
2  6
4  7
sage: Q.pp()
1  3  5
2  4
6  7
```

The inverse RSK algorithm is implemented as follows:

```
sage: RSK_inverse(P,Q, output='permutation')
[4, 2, 7, 3, 6, 1, 5]
```

We can verify that the RSK algorithm is a bijection:

12.1.2 Tutorial: Using the Sage notebook, navigating the help system, first exercises

This worksheet is based on William Stein's JPL09__intro_to_sage.sws worksheet and the Sage days 20.5_demo worksheet and aims to be an interactive introduction to Sage through exercises. You will learn how to use the notebook and call the help.

Making this help page into a worksheet

If you are browsing this document as a static web page, you can see all the examples; however you need to copy-paste them one by one to experiment with them. Use the Upload worksheet button of the notebook and copy-paste the URL of this page to obtain an editable copy in your notebook.

If you are browsing this document as part of Sage's live documentation, you can play with the examples directly here; however your changes will be lost when you close this page. Use Copy worksheet from the File... menu at the top of this page to get an editable copy in your notebook.

Both in the live tutorial and in the notebook, you can clear all output by selecting Delete All Output from the Action... menu next to the File... menu at the top of the worksheet.

Entering, Editing and Evaluating Input

To evaluate code in the Sage Notebook, type the code into an input cell and press shift-enter or click the evaluate link. Try it now with a simple expression (e.g., 2+3). The first time you evaluate a cell takes longer than subsequent times since a new Sage process is started:

```
sage: 2 + 3
5
sage: # edit here
sage: # edit here
```

To create *new input cells*, click the blue line that appears between cells when you move your mouse around. Try it now:

```
sage: 1 + 1
2
sage: # edit here
```

You can go back and edit any cell by clicking in it (or using the arrow keys on your keyboard to move up or down). Go back and change your 2 + 3 above to 3 + 3 and re-evaluate it. An empty cell can be deleted with backspace.

You can also *edit this text* right here by double clicking on it, which will bring up the TinyMCE Javascript text editor. You can even put embedded mathematics like this \$sin(x) - y^3\$ by using dollar signs just like in TeX or LaTeX.

Help systems

There are various ways of getting help in Sage.

- navigate through the documentation (there is a link Help at the top right of the worksheet),
- tab completion,
- · contextual help.

We detail below the latter two methods through examples.

Completion and contextual documentation

Start typing something and press the tab key. The interface tries to complete it with a command name. If there is more than one completion, then they are all presented to you. Remember that Sage is case sensitive, i.e. it differentiates upper case from lower case. Hence the tab completion of klein won't show you the KleinFourGroup command that builds the group $\mathbf{Z}/2 \times \mathbf{Z}/2$ as a permutation group. Try it on the next cells:

```
sage: klein<tab>
sage: Klein<tab>
```

To see documentation and examples for a command, type a question mark? at the end of the command name and press the tab key as in:

```
sage: KleinFourGroup?<tab>
sage: # edit here
```

Exercise A What is the largest prime factor of 600851475143? sage: factor?<tab> sage: # edit here

In the above manipulations we have not stored any data for later use. This can be done in Sage with the = symbol as in:

```
sage: a = 3
sage: b = 2
sage: print a+b
5
```

This can be understood as Sage evaluating the expression to the right of the = sign and creating the appropriate object, and then associating that object with a label, given by the left-hand side (see the foreword of *Tutorial: Objects and Classes in Python and Sage* for details). Multiple assignments can be done at once:

```
sage: a,b = 2,3
sage: print a,b
2 3
```

This allows us to swap the values of two variables directly:

```
sage: a,b = 2,3
sage: a,b = b,a
sage: print a,b
3 2
```

We can also assign a common value to several variables simultaneously:

```
sage: c = d = 1
sage: c, d
(1, 1)
sage: d = 2
sage: c, d
(1, 2)
```

Note that when we use the word *variable* in the computer-science sense we mean "a label attached to some data stored by Sage". Once an object is created, some *methods* apply to it. This means *functions* but instead of writing **f(my_object)** you write **my_object.f()**:

```
sage: p = 17
sage: p.is_prime()
True
```

See *Tutorial: Objects and Classes in Python and Sage* for details. To know all methods of an object you can once more use tab-completion. Write the name of the object followed by a dot and then press tab:

```
sage: a.<tab>
sage: # edit here
```

Exercise B

Create the permutation 51324 and assign it to the variable p.

```
sage: Permutation?<tab>
```

sage: # edit here

What is the inverse of p?

```
sage: p.inv<tab>
sage: # edit here
```

Does p have the pattern 123? What about 1234? And 312? (even if you don't know what a pattern is, you should be able to find a command that does this).

```
sage: p.pat<tab>
sage: # edit here
```

Some linear algebra

Exercise C

Use the matrix () command to create the following matrix.

$$M = \left(\begin{array}{cccc} 10 & 4 & 1 & 1\\ 4 & 6 & 5 & 1\\ 1 & 5 & 6 & 4\\ 1 & 1 & 4 & 10 \end{array}\right)$$

sage: matrix?<tab>

sage: # edit here

Then, using methods of the matrix,

- 1. Compute the determinant of the matrix.
- 2. Compute the echelon form of the matrix.
- 3. Compute the eigenvalues of the matrix.
- 4. Compute the kernel of the matrix.
- 5. Compute the LLL decomposition of the matrix (and lookup the documentation for what LLL is if needed!)

```
sage: # edit here
sage: # edit here
```

Now that you know how to access the different methods of matrices,

- 6. Create the vector v = (1, -1, -1, 1).
- 7. Compute the two products: $M \cdot v$ and $v \cdot M$. What mathematically borderline operation is Sage doing implicitly?

sage: vector?<tab>

sage: # edit here

Note: Vectors in Sage are row vectors. A method such as eigenspaces might not return what you expect, so it is best to specify eigenspaces_left or eigenspaces_right instead. Same thing for kernel (left_kernel or right_kernel), and so on.

Some Plotting

The plot () command allows you to draw plots of functions. Recall that you can access the documentation by pressing the tab key after writing plot? in a cell:

```
sage: plot?<tab>
sage: # edit here
```

Here is a simple example:

```
sage: var('x')  # make sure x is a symbolic variable
x
sage: plot(\sin(x^2), (x,0,10))
Graphics object consisting of 1 graphics primitive
```

Here is a more complicated plot. Try to change every single input to the plot command in some way, evaluating to see what happens:

```
sage: P = plot(sin(x^2), (x, -2, 2), rgbcolor=(0.8, 0, 0.2), thickness=3, linestyle='--', fill='axis') sage: show(P, gridlines=True)
```

Above we used the show () command to show a plot after it was created. You can also use P. show instead:

```
sage: P.show(gridlines=True)
```

Try putting the cursor right after P. show (and pressing tab to get a list of the options for how you can change the values of the given inputs.

```
sage: P.show(
```

Plotting multiple functions at once is as easy as adding them together:

```
sage: P1 = plot(sin(x), (x,0,2*pi))
sage: P2 = plot(cos(x), (x,0,2*pi), rgbcolor='red')
sage: P1 + P2
Graphics object consisting of 2 graphics primitives
```

Symbolic Expressions

Here is an example of a symbolic function:

```
sage: f(x) = x^4 - 8*x^2 - 3*x + 2
sage: f(x)
x^4 - 8*x^2 - 3*x + 2
sage: f(-3)
20
```

This is an example of a function in the *mathematical* variable x. When Sage starts, it defines the symbol x to be a mathematical variable. If you want to use other symbols for variables, you must define them first:

```
sage: x^2
x^2
sage: u + v
Traceback (most recent call last):
...
NameError: name 'u' is not defined
sage: var('u v')
(u, v)
sage: u + v
u + v
```

Still, it is possible to define symbolic functions without first defining their variables:

```
sage: f(w) = w^2
sage: f(3)
9
```

In this case those variables are defined implicitly:

```
sage: W
```

Exercise D

Define the symbolic function $f(x) = x \sin(x^2)$. Plot f on the domain [-3,3] and color it red. Use the find_root () method to numerically approximate the root of f on the interval [1,2]:

```
sage: # edit here
```

Compute the tangent line to f at x = 1:

```
sage: # edit here
```

Plot f and the tangent line to f at x = 1 in one image:

sage: # edit here

Exercise E (Advanced)

Solve the following equation for y:

$$y = 1 + xy^2$$

There are two solutions, take the one for which $\lim_{x\to 0} y(x) = 1$. (Don't forget to create the variables x and y!).

```
sage: # edit here
```

Expand y as a truncated Taylor series around 0 and containing n = 10 terms.

```
sage: # edit here
```

Do you recognize the coefficients of the Taylor series expansion? You might want to use the On-Line Encyclopedia of Integer Sequences, or better yet, Sage's class OEIS which queries the encyclopedia:

```
sage: oeis?<tab>
```

sage: # edit here

Congratulations for completing your first Sage tutorial!

12.1.3 Abelian Sandpile Model

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Introduction

These notes provide an introduction to Dhar's abelian sandpile model (ASM) and to Sage Sandpiles, a collection of tools in Sage for doing sandpile calculations. For a more thorough introduction to the theory of the ASM, the papers *Chip-Firing and Rotor-Routing on Directed Graphs* [H], by Holroyd et al. and *Riemann-Roch and Abel-Jacobi Theory on a Finite Graph* by Baker and Norine [BN] are recommended.

To describe the ASM, we start with a *sandpile graph*: a directed multigraph Γ with a vertex s that is accessible from every vertex (except possibly s, itself). By *multigraph*, we mean that each edge of Γ is assigned a nonnegative integer

weight. To say s is accessible from some vertex v means that there is a sequence of directed edges starting at v and ending at s. We call s the sink of the sandpile graph, even though it might have outgoing edges, for reasons that will be made clear in a moment.

We denoted the vertices of Γ by V and define $\tilde{V} = V \setminus \{s\}$.

Configurations and divisors

A configuration on Γ is an element of $\mathbb{N}\tilde{V}$, i.e., the assignment of a nonnegative integer to each nonsink vertex. We think of each integer as a number of grains of sand being placed at the corresponding vertex. A divisor on Γ is an element of $\mathbb{Z}V$, i.e., an element in the free abelian group on all of the vertices. In the context of divisors, it is sometimes useful to think of assigning dollars to each vertex, with negative integers signifying a debt.

Stabilization

A configuration c is stable at a vertex $v \in \tilde{V}$ if c(v) < out-degree(v), and c itself is stable if it is stable at each nonsink vertex. Otherwise, c is unstable. If c is unstable at v, the vertex v can be fired (toppled) by removing out-degree(v) grains of sand from v and adding grains of sand to the neighbors of sand, determined by the weights of the edges leaving v.

Despite our best intentions, we sometimes consider firing a stable vertex, resulting in a configuration with a "negative amount" of sand at that vertex. We may also *reverse-firing* a vertex, absorbing sand from the vertex's neighbors.

Example. Consider the graph:

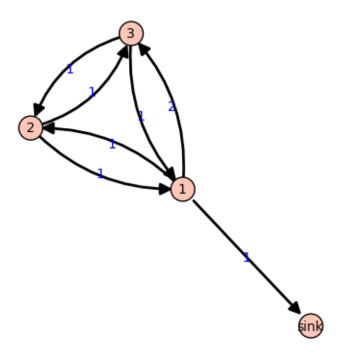


Figure 12.1: Γ

All edges have weight 1 except for the edge from vertex 1 to vertex 3, which has weight 2. If we let c = (5, 0, 1) with

the indicated number of grains of sand on vertices 1, 2, and 3, respectively, then only vertex 1, whose out-degree is 4, is unstable. Firing vertex 1 gives a new configuration c'=(1,1,3). Here, 4 grains have left vertex 1. One of these has gone to the sink vertex (and forgotten), one has gone to vertex 1, and two have gone to vertex 2, since the edge from 1 to 2 has weight 2. Vertex 3 in the new configuration is now unstable. The Sage code for this example follows.

```
sage: g = {'sink':{},
         1:{'sink':1, 2:1, 3:2},
         2:{1:1, 3:1},
. . . . :
          3:{1:1, 2:1}}
. . . . :
sage: S = Sandpile(g, 'sink') # create the sandpile
sage: S.show(edge_labels=true) # display the graph
Create the configuration:
sage: c = SandpileConfig(S, \{1:5, 2:0, 3:1\})
sage: S.out_degree()
{1: 4, 2: 2, 3: 2, 'sink': 0}
Fire vertex one:
sage: c.fire_vertex(1)
{1: 1, 2: 1, 3: 3}
The configuration is unchanged:
sage: c
{1: 5, 2: 0, 3: 1}
Repeatedly fire vertices until the configuration becomes stable:
sage: c.stabilize()
{1: 2, 2: 1, 3: 1}
Alternatives:
sage: ~c
                      # shorthand for c.stabilize()
{1: 2, 2: 1, 3: 1}
sage: c.stabilize(with_firing_vector=true)
[\{1: 2, 2: 1, 3: 1\}, \{1: 2, 2: 2, 3: 3\}]
```

Since vertex 3 has become unstable after firing vertex 1, it can be fired, which causes vertex 2 to become unstable, etc. Repeated firings eventually lead to a stable configuration. The last line of the Sage code, above, is a list, the first element of which is the resulting stable configuration, (2,1,1). The second component records how many times each vertex fired in the stabilization.

Since the sink is accessible from each nonsink vertex and never fires, every configuration will stabilize after a finite number of vertex-firings. It is not obvious, but the resulting stabilization is independent of the order in which unstable vertices are fired. Thus, each configuration stabilizes to a unique stable configuration.

Laplacian

Fix an order on the vertices of Γ . The *Laplacian* of Γ is

$$L := D - A$$

where D is the diagonal matrix of out-degrees of the vertices and A is the adjacency matrix whose (i,j)-th entry is the weight of the edge from vertex i to vertex j, which we take to be 0 if there is no edge. The *reduced Laplacian*, \tilde{L} , is the submatrix of the Laplacian formed by removing the row and column corresponding to the sink vertex. Firing a vertex of a configuration is the same as subtracting the corresponding row of the reduced Laplacian.

Example. (Continued.)

```
sage: S.vertices() # the ordering of the vertices
[1, 2, 3, 'sink']
sage: S.laplacian()
[4 -1 -2 -1]
[-1 \ 2 \ -1 \ 0]
[-1 -1 2
          01
[0 0 0 0]
sage: S.reduced_laplacian()
[ 4 -1 -2]
[-1 \ 2 \ -1]
[-1 \ -1 \ 2]
The configuration we considered previously:
sage: c = SandpileConfig(S, [5,0,1])
sage: c
{1: 5, 2: 0, 3: 1}
Firing vertex 1 is the same as subtracting the
corresponding row from the reduced Laplacian:
sage: c.fire_vertex(1).values()
[1, 1, 3]
sage: S.reduced_laplacian()[0]
(4, -1, -2)
sage: vector([5,0,1]) - vector([4,-1,-2])
(1, 1, 3)
```

Recurrent elements

Imagine an experiment in which grains of sand are dropped one-at-a-time onto a graph, pausing to allow the configuration to stabilize between drops. Some configurations will only be seen once in this process. For example, for most graphs, once sand is dropped on the graph, no addition of sand+stabilization will result in a graph empty of sand. Other configurations—the so-called *recurrent configurations*—will be seen infinitely often as the process is repeated indefinitely.

To be precise, a configuration c is *recurrent* if (i) it is stable, and (ii) given any configuration a, there is a configuration b such that $c = \operatorname{stab}(a+b)$, the stabilization of a+b.

The maximal-stable configuration, denoted c_{\max} is defined by $c_{\max}(v) = \text{out-degree}(v) - 1$ for all nonsink vertices v. It is clear that c_{\max} is recurrent. Further, it is not hard to see that a configuration is recurrent if and only if it has the form $\text{stab}(a + c_{\max})$ for some configuration a.

Example. (Continued.)

```
sage: S.recurrents(verbose=false)
[[3, 1, 1], [2, 1, 1], [3, 1, 0]]
sage: c = SandpileConfig(S, [2,1,1])
sage: c
{1: 2, 2: 1, 3: 1}
sage: c.is_recurrent()
```

```
sage: S.max_stable()
{1: 3, 2: 1, 3: 1}
Adding any configuration to the max-stable configuration and stabilizing
yields a recurrent configuration.
sage: x = SandpileConfig(S, [1,0,0])
sage: x + S.max_stable()
{1: 4, 2: 1, 3: 1}
Use & to add and stabilize:
sage: c = x & S.max_stable()
sage: c
{1: 3, 2: 1, 3: 0}
sage: c.is_recurrent()
True
Note the various ways of performing addition + stabilization:
sage: m = S.max_stable()
sage: (x + m).stabilize() == ~(x + m)
True
sage: (x + m).stabilize() == x & m
True
```

Burning Configuration

A burning configuration is a nonnegative integer-linear combination of the rows of the reduced Laplacian matrix having nonnegative entries and such that every vertex has a path from some vertex in its support. The corresponding burning script gives the integer-linear combination needed to obtain the burning configuration. So if b is the burning configuration, σ is its script, and \tilde{L} is the reduced Laplacian, then $\sigma \tilde{L} = b$. The minimal burning configuration is the one with the minimal script (its components are no larger than the components of any other script for a burning configuration).

The following are equivalent for a configuration c with burning configuration b having script σ :

- c is recurrent;
- c + b stabilizes to c;
- the firing vector for the stabilization of c + b is σ .

The burning configuration and script are computed using a modified version of Speer's script algorithm. This is a generalization to directed multigraphs of Dhar's burning algorithm.

Example.

```
sage: g = {0:{},1:{0:1,3:1,4:1},2:{0:1,3:1,5:1},
...: 3:{2:1,5:1},4:{1:1,3:1},5:{2:1,3:1}}
sage: G = Sandpile(g,0)
sage: G.burning_config()
{1: 2, 2: 0, 3: 1, 4: 1, 5: 0}
sage: G.burning_config().values()
[2, 0, 1, 1, 0]
sage: G.burning_script()
{1: 1, 2: 3, 3: 5, 4: 1, 5: 4}
```

```
sage: G.burning_script().values()
[1, 3, 5, 1, 4]
sage: matrix(G.burning_script().values())*G.reduced_laplacian()
[2 0 1 1 0]
```

Sandpile group

The collection of stable configurations forms a commutative monoid with addition defined as ordinary addition followed by stabilization. The identity element is the all-zero configuration. This monoid is a group exactly when the underlying graph is a DAG (directed acyclic graph).

The recurrent elements form a submonoid which turns out to be a group. This group is called the *sandpile group* for Γ , denoted $S(\Gamma)$. Its identity element is usually not the all-zero configuration (again, only in the case that Γ is a DAG). So finding the identity element is an interesting problem.

Let n = |V| - 1 and fix an ordering of the nonsink vertices. Let $\tilde{\mathcal{L}} \subset \mathbb{Z}^n$ denote the column-span of \tilde{L}^t , the transpose of the reduced Laplacian. It is a theorem that

$$\mathcal{S}(\Gamma) \approx \mathbb{Z}^n/\tilde{\mathcal{L}}.$$

Thus, the number of elements of the sandpile group is $\det \tilde{L}$, which by the matrix-tree theorem is the number of weighted trees directed into the sink.

Example. (Continued.)

Self-organized criticality

The sandpile model was introduced by Bak, Tang, and Wiesenfeld in the paper, Self-organized criticality: an explanation of 1/f noise [BTW]. The term self-organized criticality has no precise definition, but can be loosely taken to describe a system that naturally evolves to a state that is barely stable and such that the instabilities are described by a power law. In practice, self-organized criticality is often taken to mean like the sandpile model on a grid-graph. The grid graph is just a grid with an extra sink vertex. The vertices on the interior of each side have one edge to the sink, and the corner vertices have an edge of weight 2. Thus, every nonsink vertex has out-degree 4.

Imagine repeatedly dropping grains of sand on and empty grid graph, allowing the sandpile to stabilize in between. At first there is little activity, but as time goes on, the size and extent of the avalanche caused by a single grain of sand becomes hard to predict. Computer experiments—I do not think there is a proof, yet—indicate that the distribution of avalanche sizes obeys a power law with exponent -1. In the example below, the size of an avalanche is taken to be the sum of the number of times each vertex fires.

Example (distribution of avalanche sizes).

```
sage: S = sandpiles.Grid(10,10)
sage: m = S.max_stable()
sage: a = []
sage: for i in range(10000): # long time (15s on sage.math, 2012)
....: m = m.add_random()
....: m, f = m.stabilize(true)
....: a.append(sum(f.values()))
....
sage: p = list_plot([[log(i+1),log(a.count(i))] for i in [0..max(a)] if a.count(i)]) # long time
sage: p.axes_labels(['log(N)','log(D(N))']) # long time
sage: p # long time
Graphics object consisting of 1 graphics primitive
```

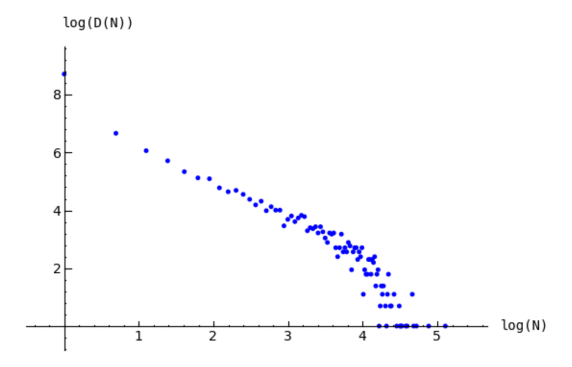


Figure 12.2: Distribution of avalanche sizes

Note: In the above code, m.stabilize(true) returns a list consisting of the stabilized configuration and the firing vector. (Omitting true would give just the stabilized configuration.)

Divisors and Discrete Riemann surfaces

A reference for this section is Riemann-Roch and Abel-Jacobi theory on a finite graph [BN].

A divisor on Γ is an element of the free abelian group on its vertices, including the sink. Suppose, as above, that the n+1 vertices of Γ have been ordered, and that $\mathcal L$ is the column span of the transpose of the Laplacian. A divisor is

then identified with an element $D \in \mathbb{Z}^{n+1}$ and two divisors are *linearly equivalent* if they differ by an element of \mathcal{L} . A divisor E is *effective*, written $E \geq 0$, if $E(v) \geq 0$ for each $v \in V$, i.e., if $E \in \mathbb{N}^{n+1}$. The *degree* of a divisor, D, is $deg(D) := \sum_{v \in V} D(v)$. The divisors of degree zero modulo linear equivalence form the *Picard group*, or *Jacobian* of the graph. For an undirected graph, the Picard group is isomorphic to the sandpile group.

The complete linear system for a divisor D, denoted |D|, is the collection of effective divisors linearly equivalent to D.

Riemann-Roch To describe the Riemann-Roch theorem in this context, suppose that Γ is an undirected, unweighted graph. The *dimension*, r(D) of the linear system |D| is -1 if $|D| = \emptyset$ and otherwise is the greatest integer s such that $|D-E| \neq 0$ for all effective divisors E of degree s. Define the *canonical divisor* by $K = \sum_{v \in V} (\deg(v) - 2)v$ and the *genus* by g = #(E) - #(V) + 1. The Riemann-Roch theorem says that for any divisor D,

$$r(D) - r(K - D) = \deg(D) + 1 - g.$$

Example.:

```
sage: G = sandpiles.Complete(5) # the sandpile on the complete graph with 5 vertices
A divisor on the graph:
sage: D = SandpileDivisor(G, [1,2,2,0,2])
Verify the Riemann-Roch theorem:
sage: K = G.canonical_divisor()
sage: D.rank() - (K - D).rank() == D.deg() + 1 - G.genus()
True
The effective divisors linearly equivalent to D:
sage: D.effective_div(False)
[[0, 1, 1, 4, 1], [1, 2, 2, 0, 2], [4, 0, 0, 3, 0]]
The nonspecial divisors up to linear equivalence (divisors of degree
g-1 with empty linear systems)
sage: N = G.nonspecial_divisors()
sage: [E.values() for E in N[:5]]
                                    # the first few
[[-1, 0, 1, 2, 3],
[-1, 0, 1, 3, 2],
[-1, 0, 2, 1, 3],
[-1, 0, 2, 3, 1],
[-1, 0, 3, 1, 2]
sage: len(N)
sage: len(N) == G.h_vector()[-1]
True
```

Picturing linear systems Fix a divisor D. There are at least two natural graphs associated with linear system associated with D. First, consider the directed graph with vertex set |D| and with an edge from vertex E to vertex F if F is attained from E by firing a single unstable vertex.

```
sage: S = Sandpile(graphs.CycleGraph(6),0)
sage: D = SandpileDivisor(S, [1,1,1,1,2,0])
sage: D.is_alive()
True
```

```
sage: eff = D.effective_div()
sage: firing_graph(S,eff).show3d(edge_size=.005,vertex_size=0.01,iterations=500)
```

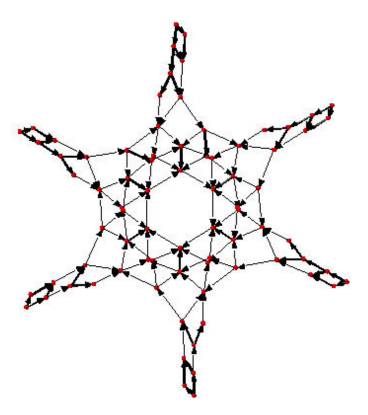


Figure 12.3: Complete linear system for (1,1,1,1,2,0) on C_6 : single firings

The second graph has the same set of vertices but with an edge from E to F if F is obtained from E by firing all unstable vertices of E.

```
sage: S = Sandpile(graphs.CycleGraph(6),0)
sage: D = SandpileDivisor(S, [1,1,1,1,2,0])
sage: eff = D.effective_div()
sage: parallel_firing_graph(S,eff).show3d(edge_size=.005,vertex_size=0.01,iterations=500)
```

Note that in each of the examples, above, starting at any divisor in the linear system and following edges, one is eventually led into a cycle of length 6 (cycling the divisor (1,1,1,1,2,0)). Thus, D.alive() returns True. In Sage, one would be able to rotate the above figures to get a better idea of the structure.

Algebraic geometry of sandpiles

Affine Let n=|V|-1, and fix an ordering on the nonsink vertices of Γ . let $\tilde{\mathcal{L}}\subset\mathbb{Z}^n$ denote the column-span of $\tilde{\mathcal{L}}^t$, the transpose of the reduced Laplacian. Label vertex i with the indeterminate x_i , and let $\mathbb{C}[\Gamma_s]=\mathbb{C}[x_1,\ldots,x_n]$. (Here, s denotes the sink vertex of Γ .) The *sandpile ideal* or *toppling ideal*, first studied by Cori, Rossin, and Salvy [CRS] for undirected graphs, is the lattice ideal for $\tilde{\mathcal{L}}$:

$$I = I(\Gamma_s) := \{ x^u - x^v : u - v \in \tilde{\mathcal{L}} \} \subset \mathbb{C}[\Gamma_s],$$

where $x^u := \prod_{i=1}^n x^{u_i}$ for $u \in \mathbb{Z}^n$.

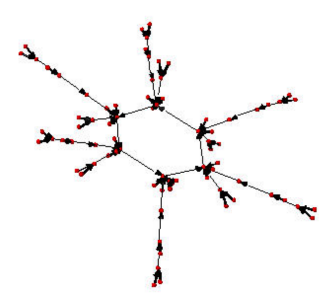


Figure 12.4: Complete linear system for (1,1,1,1,2,0) on C_6 : parallel firings

For each $c \in \mathbb{Z}^n$ define $t(c) = x^{c^+} - x^{c^-}$ where $c_i^+ = \max\{c_i, 0\}$ and $c^- = \max\{-c_i, 0\}$ so that $c = c^+ - c^-$. Then, for each $\sigma \in \mathbb{Z}^n$, define $T(\sigma) = t(\tilde{L}^t\sigma)$. It then turns out that

$$I = (T(e_1), \dots, T(e_n), x^b - 1)$$

where e_i is the *i*-th standard basis vector and b is any burning configuration.

The affine coordinate ring, $\mathbb{C}[\Gamma_s]/I$, is isomorphic to the group algebra of the sandpile group, $\mathbb{C}[\mathcal{S}(\Gamma)]$.

The standard term-ordering on $\mathbb{C}[\Gamma_s]$ is graded reverse lexigraphical order with $x_i > x_j$ if vertex v_i is further from the sink than vertex v_j . (There are choices to be made for vertices equidistant from the sink). If σ_b is the script for a burning configuration (not necessarily minimal), then

$$\{T(\sigma): \sigma \leq \sigma_b\}$$

is a Groebner basis for I.

Projective Now let $\mathbb{C}[\Gamma] = \mathbb{C}[x_0, x_1, \dots, x_n]$, where x_0 corresponds to the sink vertex. The *homogeneous sandpile ideal*, denoted I^h , is obtaining by homogenizing I with respect to x_0 . Let L be the (full) Laplacian, and $L \subset \mathbb{Z}^{n+1}$ be the column span of its transpose, L^t . Then I^h is the lattice ideal for L:

$$I^h = I^h(\Gamma) := \{x^u - x^v : u - v \in \mathcal{L}\} \subset \mathbb{C}[\Gamma].$$

This ideal can be calculated by saturating the ideal

$$(T(e_i): i = 0, \dots n)$$

with respect to the product of the indeterminates: $\prod_{i=0}^{n} x_i$ (extending the T operator in the obvious way). A Groebner basis with respect to the degree lexicographic order describe above (with x_0 the smallest vertex), is obtained by homogenizing each element of the Groebner basis for the non-homogeneous sandpile ideal with respect to x_0 .

Example.

```
sage: g = \{0:\{\}, 1:\{0:1, 3:1, 4:1\}, 2:\{0:1, 3:1, 5:1\},
         3:\{2:1,5:1\},4:\{1:1,3:1\},5:\{2:1,3:1\}\}
sage: S = Sandpile(g, 0)
sage: S.ring()
Multivariate Polynomial Ring in x5, x4, x3, x2, x1, x0 over Rational Field
The homogeneous sandpile ideal:
sage: S.ideal()
Ideal (x2 - x0, x3^2 - x5*x0, x5*x3 - x0^2, x4^2 - x3*x1, x5^2 - x3*x0,
x1^3 - x4*x3*x0, x4*x1^2 - x5*x0^2) of Multivariate Polynomial Ring
in x5, x4, x3, x2, x1, x0 over Rational Field
The generators of the ideal:
sage: S.ideal(true)
[x2 - x0,
x3^2 - x5*x0,
x5*x3 - x0^2,
x4^2 - x3*x1,
x5^2 - x3*x0,
x1^3 - x4*x3*x0,
x4*x1^2 - x5*x0^2
Its resolution:
sage: S.resolution() # long time
'R^1 <-- R^7 <-- R^19 <-- R^25 <-- R^16 <-- R^4'
and Betti table:
sage: S.betti() # long time
         0 1 2 3
                                        5
             1
    0: 1
                4
                      6
                            2
    1:
                      7
                           7
    2:
                2
                                  2
                     6
                          16
                                 14
    3:
                                        4
total: 1 7 19 25 16 4
The Hilbert function:
sage: S.hilbert_function()
[1, 5, 11, 15]
and its first differences (which counts the number of superstable
configurations in each degree):
sage: S.h_vector()
[1, 4, 6, 4]
sage: x = [i.deg() for i in S.superstables()]
sage: sorted(x)
```

```
[0, 1, 1, 1, 1, 2, 2, 2, 2, 2, 2, 3, 3, 3, 3]
```

The degree in which the Hilbert function equals the Hilbert polynomial, the latter always being a constant in the case of a sandpile ideal:

```
sage: S.postulation()
```

Zeros The *zero set* for the sandpile ideal *I* is

$$Z(I)=\{p\in\mathbb{C}^n: f(p)=0 \text{ for all } f\in I\},$$

the set of simultaneous zeros of the polynomials in I. Letting S^1 denote the unit circle in the complex plane, Z(I) is a finite subgroup of $S^1 \times \cdots \times S^1 \subset \mathbb{C}^n$, isomorphic to the sandpile group. The zero set is actually linearly isomorphic to a faithful representation of the sandpile group on \mathbb{C}^n .

Example. (Continued.)

```
sage: S = Sandpile(\{0: \{\}, 1: \{2: 2\}, 2: \{0: 4, 1: 1\}\}, 0)
sage: S.ideal().gens()
[x1^2 - x2^2, x1*x2^3 - x0^4, x2^5 - x1*x0^4]
Approximation to the zero set (setting 'x_0 = 1''):
sage: S.solve()
[[-0.707107 + 0.707107 \times I, 0.707107 - 0.707107 \times I],
[-0.707107 - 0.707107*I, 0.707107 + 0.707107*I],
[-I, -I],
[I, I],
[0.707107 + 0.707107 \times I, -0.707107 - 0.707107 \times I],
[0.707107 - 0.707107 \times I, -0.707107 + 0.707107 \times I],
[1, 1],
[-1, -1]
sage: len(_) == S.group_order()
True
The zeros are generated as a group by a single vector:
sage: S.points()
[[e^{(1/4*I*pi)}, e^{(-3/4*I*pi)}]]
```

Resolutions The homogeneous sandpile ideal, I^h , has a free resolution graded by the divisors on Γ modulo linear equivalence. (See the section on *Discrete Riemann Surfaces* for the language of divisors and linear equivalence.) Let $S = \mathbb{C}[\Gamma] = \mathbb{C}[x_0, \dots, x_n]$, as above, and let \mathfrak{S} denote the group of divisors modulo rational equivalence. Then S is graded by \mathfrak{S} by letting $\deg(x^c) = c \in \mathfrak{S}$ for each monomial x^c . The minimal free resolution of I^h has the form

$$0 \leftarrow I^h \leftarrow \bigoplus_{D \in \mathfrak{S}} S(-D)^{\beta_{0,D}} \leftarrow \bigoplus_{D \in \mathfrak{S}} S(-D)^{\beta_{1,D}} \leftarrow \cdots \leftarrow \bigoplus_{D \in \mathfrak{S}} S(-D)^{\beta_{r,D}} \leftarrow 0.$$

where the $\beta_{i,D}$ are the *Betti numbers* for I^h .

For each divisor class $D \in \mathfrak{S}$, define a simplicial complex,

$$\Delta_D := \{ I \subseteq \{0, \dots, n\} : I \subseteq \text{supp}(E) \text{ for some } E \in |D| \}.$$

The Betti number $\beta_{i,D}$ equals the dimension over \mathbb{C} of the *i*-th reduced homology group of Δ_D :

$$\beta_{i,D} = \dim_{\mathbb{C}} \tilde{H}_i(\Delta_D; \mathbb{C}).$$

```
sage: S = Sandpile({0:{},1:{}0:{}1,{}2:{}1,{}3:{}4},2:{}3:{}5},3:{}1:{}1,{}2:{}1}},0)
Representatives of all divisor classes with nontrivial homology:
sage: p = S.betti_complexes()
sage: p[0]
[\{0: -8, 1: 5, 2: 4, 3: 1\},
Simplicial complex with vertex set (1, 2, 3) and facets \{(1, 2), (3,)\}
The homology associated with the first divisor in the list:
sage: D = p[0][0]
sage: D.effective_div()
[\{0: 0, 1: 0, 2: 0, 3: 2\}, \{0: 0, 1: 1, 2: 1, 3: 0\}]
sage: [E.support() for E in D.effective_div()]
[[3], [1, 2]]
sage: D.Dcomplex()
Simplicial complex with vertex set (1, 2, 3) and facets \{(1, 2), (3,)\}
sage: D.Dcomplex().homology()
{0: Z, 1: 0}
The minimal free resolution:
sage: S.resolution()
'R^1 <-- R^5 <-- R^1'
sage: S.betti()
         0
               1
                     2 3
   0: 1
1: -
               5
                     5
total:
       1 5
                     5
sage: len(p)
11
The degrees and ranks of the homology groups for each element of the list p
(compare with the Betti table, above):
sage: [[sum(d[0].values()),d[1].betti()] for d in p]
[[2, \{0: 2, 1: 0\}],
[3, {0: 1, 1: 1, 2: 0}],
[2, {0: 2, 1: 0}],
[3, \{0: 1, 1: 1, 2: 0\}],
[2, {0: 2, 1: 0}],
[3, {0: 1, 1: 1, 2: 0}],
[2, {0: 2, 1: 0}],
[3, {0: 1, 1: 1}],
[2, \{0: 2, 1: 0\}],
[3, \{0: 1, 1: 1, 2: 0\}],
[5, {0: 1, 1: 0, 2: 1}]]
```

Complete Intersections and Arithmetically Gorenstein toppling ideals NOTE: in the previous section note that the resolution always has length n since the ideal is Cohen-Macaulay.

To do.

Betti numbers for undirected graphs To do.

Usage

Initialization

There are three main classes for sandpile structures in Sage: Sandpile, SandpileConfig, and SandpileDivisor. Initialization for Sandpile has the form

```
sage: S = Sandpile(graph, sink)
```

where graph represents a graph and sink is the key for the sink vertex. There are four possible forms for graph:

1. a Python dictionary of dictionaries:

```
sage: g = \{0: \{\}, 1: \{0: 1, 3: 1, 4: 1\}, 2: \{0: 1, 3: 1, 5: 1\}, \ldots: 3: \{2: 1, 5: 1\}, 4: \{1: 1, 3: 1\}, 5: \{2: 1, 3: 1\}\}
```

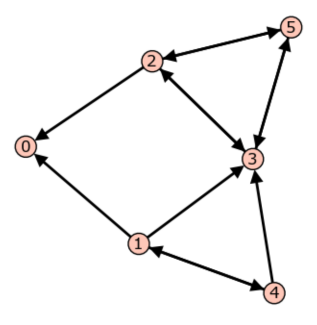


Figure 12.5: Graph from dictionary of dictionaries.

Each key is the name of a vertex. Next to each vertex name v is a dictionary consisting of pairs: vertex: weight. Each pair represents a directed edge emanating from v and ending at vertex having (non-negative integer) weight equal to weight. Loops are allowed. In the example above, all of the weights are 1.

2. a Python dictionary of lists:

```
sage: g = {0: [], 1: [0, 3, 4], 2: [0, 3, 5],
....: 3: [2, 5], 4: [1, 3], 5: [2, 3]}
```

This is a short-hand when all of the edge-weights are equal to 1. The above example is for the same displayed graph.

3. a Sage graph (of type sage.graphs.graph.Graph):

```
sage: g = graphs.CycleGraph(5)
sage: S = Sandpile(g, 0)
sage: type(g)
<class 'sage.graphs.graph.Graph'>
```

To see the types of built-in graphs, type graphs., including the period, and hit TAB.

4. a Sage digraph:

```
sage: S = Sandpile(digraphs.RandomDirectedGNC(6), 0)
sage: S.show()
```

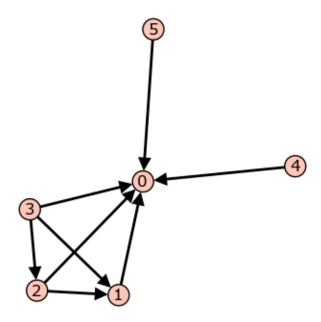


Figure 12.6: A random graph.

See sage.graphs.graph_generators for more information on the Sage graph library and graph constructors.

Each of these four formats is preprocessed by the Sandpile class so that, internally, the graph is represented by the dictionary of dictionaries format first presented. This internal format is returned by dict ():

```
sage: S = Sandpile({0:[], 1:[0, 3, 4], 2:[0, 3, 5], 3: [2, 5], 4: [1, 3], 5: [2, 3]},0)
sage: S.dict()
{0: {},
    1: {0: 1, 3: 1, 4: 1},
    2: {0: 1, 3: 1, 5: 1},
    3: {2: 1, 5: 1},
    4: {1: 1, 3: 1},
    5: {2: 1, 3: 1}}
```

Note: The user is responsible for assuring that each vertex has a directed path into the designated sink. If the sink has out-edges, these will be ignored for the purposes of sandpile calculations (but not calculations on divisors).

Code for checking whether a given vertex is a sink:

```
sage: S = Sandpile({0:[], 1:[0, 3, 4], 2:[0, 3, 5], 3: [2, 5], 4: [1, 3], 5: [2, 3]},0)
sage: [S.distance(v,0) for v in S.vertices()] # 0 is a sink
[0, 1, 1, 2, 2, 2]
sage: [S.distance(v,1) for v in S.vertices()] # 1 is not a sink
[+Infinity, 0, +Infinity, +Infinity, 1, +Infinity]
```

Methods

Here are summaries of Sandpile, SandpileConfig, and SandpileDivisor methods (functions). Each summary is followed by a list of complete descriptions of the methods. There are many more methods available for a Sandpile, e.g., those inherited from the class DiGraph. To see them all, enter dir (Sandpile) or type Sandpile., including the period, and hit TAB.

Sandpile Summary of methods.

- *all_k_config* The constant configuration with all values set to k.
- all k div The divisor with all values set to k.
- avalanche_polynomial The avalanche polynomial.
- *betti* The Betti table for the homogeneous toppling ideal.
- betti_complexes The support-complexes with non-trivial homology.
- burning_config The minimal burning configuration.
- burning_script A script for the minimal burning configuration.
- canonical_divisor The canonical divisor.
- dict A dictionary of dictionaries representing a directed graph.
- *genus* The genus: (# non-loop edges) (# vertices) + 1.
- groebner A Groebner basis for the homogeneous toppling ideal.
- group_gens A minimal list of generators for the sandpile group.
- group_order The size of the sandpile group.
- h vector The number of superstable configurations in each degree.
- *help* List of Sandpile-specific methods (not inherited from Graph).
- hilbert_function The Hilbert function of the homogeneous toppling ideal.
- *ideal* The saturated homogeneous toppling ideal.
- *identity* The identity configuration.
- in_degree The in-degree of a vertex or a list of all in-degrees.
- invariant_factors The invariant factors of the sandpile group.
- is_undirected Is the underlying graph undirected?
- jacobian_representatives Representatives for the elements of the Jacobian group.
- *laplacian* The Laplacian matrix of the graph.

- markov_chain The sandpile Markov chain for configurations or divisors.
- max_stable The maximal stable configuration.
- max_stable_div The maximal stable divisor.
- *max_superstables* The maximal superstable configurations.
- min recurrents The minimal recurrent elements.
- nonsink_vertices The nonsink vertices.
- nonspecial_divisors The nonspecial divisors.
- out_degree The out-degree of a vertex or a list of all out-degrees.
- picard_representatives Representatives of the divisor classes of degree d in the Picard group.
- points Generators for the multiplicative group of zeros of the sandpile ideal.
- postulation The postulation number of the toppling ideal.
- recurrents The recurrent configurations.
- reduced_laplacian The reduced Laplacian matrix of the graph.
- reorder_vertices A copy of the sandpile with vertex names permuted.
- resolution A minimal free resolution of the homogeneous toppling ideal.
- ring The ring containing the homogeneous toppling ideal.
- *show* Draw the underlying graph.
- *show3d* Draw the underlying graph.
- *sink* The sink vertex.
- smith_form The Smith normal form for the Laplacian.
- solve Approximations of the complex affine zeros of the sandpile ideal.
- *stable_configs* Generator for all stable configurations.
- *stationary_density* The stationary density of the sandpile.
- *superstables* The superstable configurations.
- *symmetric_recurrents* The symmetric recurrent configurations.
- *tutte_polynomial* The Tutte polynomial.
- unsaturated_ideal The unsaturated, homogeneous toppling ideal.
- version The version number of Sage Sandpiles.
- zero_config The all-zero configuration.
- *zero_div* The all-zero divisor.

Complete descriptions of Sandpile methods.

— all_k_config(k)

The constant configuration with all values set to k.

INPUT:

k - integer

OUTPUT:

SandpileConfig

EXAMPLES:

```
sage: s = sandpiles.Diamond()
sage: s.all_k_config(7)
{1: 7, 2: 7, 3: 7}
```

— all_k_div(k)

The divisor with all values set to k.

INPUT:

k - integer

OUTPUT:

SandpileDivisor

EXAMPLES:

```
sage: S = sandpiles.House()
sage: S.all_k_div(7)
{0: 7, 1: 7, 2: 7, 3: 7, 4: 7}
```

— avalanche_polynomial(multivariable=True)

The avalanche polynomial. See NOTE for details.

INPUT:

multivariable - (default: True) boolean

OUTPUT:

polynomial

EXAMPLES:

```
sage: s = sandpiles.Complete(4)
sage: s.avalanche_polynomial()
9*x0*x1*x2 + 2*x0*x1 + 2*x0*x2 + 2*x1*x2 + 3*x0 + 3*x1 + 3*x2 + 24
sage: s.avalanche_polynomial(False)
9*x0^3 + 6*x0^2 + 9*x0 + 24
```

Note: For each nonsink vertex v, let x_v be an indeterminate. If (r,v) is a pair consisting of a recurrent r and nonsink vertex v, then for each nonsink vertex w, let n_w be the number of times vertex w fires in the stabilization of r+v. Let M(r,v) be the monomial $\prod_w x_w^{n_w}$, i.e., the exponent records the vector of n_w as w ranges over the nonsink vertices. The avalanche polynomial is then the sum of M(r,v) as r ranges over the recurrents and v ranges over the nonsink vertices. If multivariable is False, then set all the indeterminates equal to each other (and, thus, only count the number of vertex firings in the stabilizations, forgetting which particular vertices fired).

— betti(verbose=True)

The Betti table for the homogeneous toppling ideal. If verbose is True, it prints the standard Betti table, otherwise, it returns a less formated table.

INPUT:

```
verbose - (default: True) boolean
```

OUTPUT:

Betti numbers for the sandpile

EXAMPLES:

```
sage: S = sandpiles.Diamond()
sage: S.betti()
                 2
             1
                    3
       0
       1
   1:
            2
   2:
                      4
            4
total: 1 6
                 9
                      4
sage: S.betti(False)
[1, 6, 9, 4]
```

— betti_complexes()

The support-complexes with non-trivial homology. (See NOTE.)

OUTPUT:

list (of pairs [divisors, corresponding simplicial complex])

EXAMPLES:

```
sage: S = Sandpile({0:{},1:{}0:{}1,{}2:{}1,{}3:{}4},2:{}3:{}5},3:{}1:{}1,{}2:{}1}},0)
sage: p = S.betti_complexes()
sage: p[0]
[\{0: -8, 1: 5, 2: 4, 3: 1\},  Simplicial complex with vertex set (1, 2, 3) and facets \{(1, 2), (3,)\}
sage: S.resolution()
'R^1 <-- R^5 <-- R^5 <-- R^1'
sage: S.betti()
             1
   0: 1 -
   1:
                5
   2:
                            1
total: 1 5 5 1
sage: len(p)
11
sage: p[0][1].homology()
{0: Z, 1: 0}
sage: p[-1][1].homology()
{0: 0, 1: 0, 2: Z}
```

Note: A support-complex is the simplicial complex formed from the supports of the divisors in a linear system.

— burning_config()

The minimal burning configuration.

OUTPUT:

dict (configuration)

Note: The burning configuration and script are computed using a modified version of Speer's script algorithm. This is a generalization to directed multigraphs of Dhar's burning algorithm.

A burning configuration is a nonnegative integer-linear combination of the rows of the reduced Laplacian matrix having nonnegative entries and such that every vertex has a path from some vertex in its support. The corresponding burning script gives the integer-linear combination needed to obtain the burning configuration. So if b is the burning configuration, σ is its script, and \tilde{L} is the reduced Laplacian, then $\sigma \cdot \tilde{L} = b$. The minimal burning configuration is the one with the minimal script (its components are no larger than the components of any other script for a burning configuration).

The following are equivalent for a configuration c with burning configuration b having script σ :

- c is recurrent;
- c + b stabilizes to c;
- the firing vector for the stabilization of c + b is σ .

— burning script()

A script for the minimal burning configuration.

OUTPUT:

dict

EXAMPLES:

```
sage: g = {0:{},1:{0:1,3:1,4:1},2:{0:1,3:1,5:1},\
3:{2:1,5:1},4:{1:1,3:1},5:{2:1,3:1}}
sage: S = Sandpile(g,0)
sage: S.burning_config()
{1: 2, 2: 0, 3: 1, 4: 1, 5: 0}
sage: S.burning_config().values()
[2, 0, 1, 1, 0]
sage: S.burning_script()
{1: 1, 2: 3, 3: 5, 4: 1, 5: 4}
sage: script = S.burning_script().values()
sage: script
[1, 3, 5, 1, 4]
sage: matrix(script)*S.reduced_laplacian()
[2 0 1 1 0]
```

Note: The burning configuration and script are computed using a modified version of Speer's script algorithm. This

is a generalization to directed multigraphs of Dhar's burning algorithm.

A burning configuration is a nonnegative integer-linear combination of the rows of the reduced Laplacian matrix having nonnegative entries and such that every vertex has a path from some vertex in its support. The corresponding burning script gives the integer-linear combination needed to obtain the burning configuration. So if b is the burning configuration, s is its script, and $L_{\rm red}$ is the reduced Laplacian, then $s \cdot L_{\rm red} = b$. The minimal burning configuration is the one with the minimal script (its components are no larger than the components of any other script for a burning configuration).

The following are equivalent for a configuration c with burning configuration b having script s:

- c is recurrent;
- c + b stabilizes to c;
- the firing vector for the stabilization of c + b is s.

— canonical_divisor()

The canonical divisor. This is the divisor with deg(v) - 2 grains of sand on each vertex (not counting loops). Only for undirected graphs.

OUTPUT:

SandpileDivisor

EXAMPLES:

```
sage: S = sandpiles.Complete(4)
sage: S.canonical_divisor()
{0: 1, 1: 1, 2: 1, 3: 1}
sage: s = Sandpile({0:[1,1],1:[0,0,1,1,1]},0)
sage: s.canonical_divisor() # loops are disregarded
{0: 0, 1: 0}
```

Warning: The underlying graph must be undirected.

— dict()

A dictionary of dictionaries representing a directed graph.

OUTPUT:

dict

EXAMPLES:

```
sage: S = sandpiles.Diamond()
sage: S.dict()
{0: {1: 1, 2: 1},
    1: {0: 1, 2: 1, 3: 1},
    2: {0: 1, 1: 1, 3: 1},
    3: {1: 1, 2: 1}}
sage: S.sink()
0
```

— genus()

The genus: (# non-loop edges) - (# vertices) + 1. Only defined for undirected graphs.

OUTPUT:

integer

EXAMPLES:

```
sage: sandpiles.Complete(4).genus()
3
sage: sandpiles.Cycle(5).genus()
1
```

— groebner()

A Groebner basis for the homogeneous toppling ideal. It is computed with respect to the standard sandpile ordering (see ring).

OUTPUT:

Groebner basis

EXAMPLES:

```
sage: S = sandpiles.Diamond()
sage: S.groebner()
[x3*x2^2 - x1^2*x0, x2^3 - x3*x1*x0, x3*x1^2 - x2^2*x0, x1^3 - x3*x2*x0, x3^2 - x0^2, x2*x1 - x0^2]
```

— group_gens(verbose=True)

A minimal list of generators for the sandpile group. If verbose is False then the generators are represented as lists of integers.

INPUT:

verbose - (default: True) boolean

OUTPUT:

list of SandpileConfig (or of lists of integers if verbose is False)

EXAMPLES:

```
sage: s = sandpiles.Cycle(5)
sage: s.group_gens()
[{1: 1, 2: 1, 3: 1, 4: 0}]
sage: s.group_gens()[0].order()
5
sage: s = sandpiles.Complete(5)
sage: s.group_gens(False)
[[2, 2, 3, 2], [2, 3, 2, 2], [3, 2, 2, 2]]
sage: [i.order() for i in s.group_gens()]
[5, 5, 5]
sage: s.invariant_factors()
[1, 5, 5, 5]
```

— group_order()

The size of the sandpile group.

OUTPUT:

integer

```
sage: S = sandpiles.House()
sage: S.group_order()
11
```

— h vector()

The number of superstable configurations in each degree. Equivalently, this is the list of first differences of the Hilbert function of the (homogeneous) toppling ideal.

OUTPUT:

list of nonnegative integers

EXAMPLES:

```
sage: s = sandpiles.Grid(2,2)
sage: s.hilbert_function()
[1, 5, 15, 35, 66, 106, 146, 178, 192]
sage: s.h_vector()
[1, 4, 10, 20, 31, 40, 40, 32, 14]
```

— help(verbose=True)

List of Sandpile-specific methods (not inherited from Graph). If verbose, include short descriptions.

INPUT:

```
verbose - (default: True) boolean
```

OUTPUT:

printed string

```
sage: Sandpile.help()
For detailed help with any method FOO listed below,
enter "Sandpile.FOO?" or enter "S.FOO?" for any Sandpile S.
                      -- The constant configuration with all values set to k.
all_k_config
all_k_div
                      -- The divisor with all values set to k.
avalanche_polynomial
                      -- The avalanche polynomial.
betti
                       -- The Betti table for the homogeneous toppling ideal.
betti_complexes
                       -- The support-complexes with non-trivial homology.
burning_config
                       -- The minimal burning configuration.
burning_script
                       -- A script for the minimal burning configuration.
canonical_divisor
                      -- The canonical divisor.
                      -- A dictionary of dictionaries representing a directed graph.
dict
                      -- The genus: (# non-loop edges) - (# vertices) + 1.
genus
groebner
                      -- A Groebner basis for the homogeneous toppling ideal.
                      -- A minimal list of generators for the sandpile group.
group_gens
group_order
h_vector
help
                      -- The size of the sandpile group.
                      -- The number of superstable configurations in each degree.
                      -- List of Sandpile-specific methods (not inherited from Graph).
help
hilbert_function -- The Hilbert function of the homogeneous toppling ideal.
ideal
                      -- The saturated homogeneous toppling ideal.
                      -- The identity configuration.
identity
in_degree
                      -- The in-degree of a vertex or a list of all in-degrees.
jacobian_representatives -- Representatives for the elements of the Jacobian group.
laplacian -- The Laplacian matrix of the graph.
markov_chain
                      -- The sandpile Markov chain for configurations or divisors.
                      -- The maximal stable configuration.
max_stable
max_stable_div
                     -- The maximal stable divisor.
max_superstables
                      -- The maximal superstable configurations.
```

```
-- The minimal recurrent elements.
min_recurrents
nonsink_vertices
                         -- The nonsink vertices.
nonspecial_divisors
                         -- The nonspecial divisors.
out_degree
                          -- The out-degree of a vertex or a list of all out-degrees.
picard_representatives
                         -- Representatives of the divisor classes of degree d in the Picard group.
                          -- Generators for the multiplicative group of zeros of the sandpile ideal.
points
postulation
                          -- The postulation number of the toppling ideal.
                          -- The recurrent configurations.
recurrents
reduced_laplacian
                         -- The reduced Laplacian matrix of the graph.
reorder_vertices
                         -- A copy of the sandpile with vertex names permuted.
resolution
                         -- A minimal free resolution of the homogeneous toppling ideal.
ring
                          -- The ring containing the homogeneous toppling ideal.
show
                          -- Draw the underlying graph.
show3d
                         -- Draw the underlying graph.
sink
                         -- The sink vertex.
smith_form
                         -- The Smith normal form for the Laplacian.
solve
stable_configs
stationary_density

-- Generator for all stable configurations.

-- The stationary density of the sandpile.

-- The superstable configurations.
                         -- Approximations of the complex affine zeros of the sandpile ideal.
                         -- Generator for all stable configurations.
tutte_polynomial
                         -- The Tutte polynomial.
unsaturated_ideal
                        -- The unsaturated, homogeneous toppling ideal.
version
                         -- The version number of Sage Sandpiles.
zero_config
                         -- The all-zero configuration.
                         -- The all-zero divisor.
zero_div
```

— hilbert function()

The Hilbert function of the homogeneous toppling ideal.

OUTPUT:

list of nonnegative integers

EXAMPLES:

```
sage: s = sandpiles.Wheel(5)
sage: s.hilbert_function()
[1, 5, 15, 31, 45]
sage: s.h_vector()
[1, 4, 10, 16, 14]
```

— ideal(gens=False)

The saturated homogeneous toppling ideal. If gens is True, the generators for the ideal are returned instead.

INPUT:

```
gens - (default: False) boolean
```

OUTPUT:

ideal or, optionally, the generators of an ideal

```
sage: S = sandpiles.Diamond()
sage: S.ideal()
Ideal (x2*x1 - x0^2, x3^2 - x0^2, x1^3 - x3*x2*x0, x3*x1^2 - x2^2*x0, x2^3 - x3*x1*x0, x3*x2^2 - x1^2
sage: S.ideal(True)
[x2*x1 - x0^2, x3^2 - x0^2, x1^3 - x3*x2*x0, x3*x1^2 - x2^2*x0, x2^3 - x3*x1*x0, x3*x2^2 - x1^2*x0]
```

```
sage: S.ideal().gens() # another way to get the generators [x2*x1 - x0^2, x3^2 - x0^2, x1^3 - x3*x2*x0, x3*x1^2 - x2^2*x0, x2^3 - x3*x1*x0, x3*x2^2 - x1^2*x0]
```

— identity(verbose=True)

The identity configuration. If verbose is False, the configuration are converted to a list of integers.

INPUT:

```
verbose - (default: True) boolean
```

OUTPUT:

SandpileConfig or a list of integers If verbose is False, the configuration are converted to a list of integers.

EXAMPLES:

```
sage: s = sandpiles.Diamond()
sage: s.identity()
{1: 2, 2: 2, 3: 0}
sage: s.identity(False)
[2, 2, 0]
sage: s.identity() & s.max_stable() == s.max_stable()
True
```

- in_degree(v=None)

The in-degree of a vertex or a list of all in-degrees.

INPUT:

∨ – (optional) vertex name

OUTPUT:

integer or dict

EXAMPLES:

```
sage: s = sandpiles.House()
sage: s.in_degree()
{0: 2, 1: 2, 2: 3, 3: 3, 4: 2}
sage: s.in_degree(2)
3
```

— invariant_factors()

The invariant factors of the sandpile group.

OUTPUT:

list of integers

EXAMPLES:

```
sage: s = sandpiles.Grid(2,2)
sage: s.invariant_factors()
[1, 1, 8, 24]
```

— is undirected()

Is the underlying graph undirected? True if (u, v) is and edge if and only if (v, u) is an edge, each edge with the same weight.

OUTPUT:

boolean

EXAMPLES:

```
sage: sandpiles.Complete(4).is_undirected()
True
sage: s = Sandpile({0:[1,2], 1:[0,2], 2:[0]}, 0)
sage: s.is_undirected()
False
```

— jacobian_representatives(verbose=True)

Representatives for the elements of the Jacobian group. If verbose is False, then lists representing the divisors are returned.

INPUT:

```
verbose - (default: True) boolean
```

OUTPUT:

list of SandpileDivisor (or of lists representing divisors)

EXAMPLES:

For an undirected graph, divisors of the form $s - \deg(s) * sink as s varies over the superstables forms a distinct set of representatives for the Jacobian group.:$

```
sage: s = sandpiles.Complete(3)
sage: s.superstables(False)
[[0, 0], [0, 1], [1, 0]]
sage: s.jacobian_representatives(False)
[[0, 0, 0], [-1, 0, 1], [-1, 1, 0]]
```

If the graph is directed, the representatives described above may by equivalent modulo the rowspan of the Laplacian matrix:

```
sage: s = Sandpile({0: {1: 1, 2: 2}, 1: {0: 2, 2: 4}, 2: {0: 4, 1: 2}},0)
sage: s.group_order()
28
sage: s.jacobian_representatives()
[{0: -5, 1: 3, 2: 2}, {0: -4, 1: 3, 2: 1}]
```

Let τ be the nonnegative generator of the kernel of the transpose of the Laplacian, and let tau_s be it sink component, then the sandpile group is isomorphic to the direct sum of the cyclic group of order τ_s and the Jacobian group. In the example above, we have:

```
sage: s.laplacian().left_kernel()
Free module of degree 3 and rank 1 over Integer Ring
Echelon basis matrix:
[14 5 8]
```

Note: The Jacobian group is the set of all divisors of degree zero modulo the integer rowspan of the Laplacian matrix.

- laplacian()

The Laplacian matrix of the graph. Its rows encode the vertex firing rules.

OUTPUT:

matrix

EXAMPLES:

```
sage: G = sandpiles.Diamond()
sage: G.laplacian()
[ 2 -1 -1 0]
[-1 3 -1 -1]
[-1 -1 3 -1]
[ 0 -1 -1 2]
```

Warning: The function laplacian_matrix should be avoided. It returns the indegree version of the Laplacian.

— markov_chain(state, distrib=None)

The sandpile Markov chain for configurations or divisors. The chain starts at state. See NOTE for details.

INPUT:

- state SandpileConfig, SandpileDivisor, or list representing one of these
- distrib (optional) list of nonnegative numbers summing to 1 (representing a prob. dist.)

OUTPUT:

generator for Markov chain (see NOTE)

```
sage: s = sandpiles.Complete(4)
sage: m = s.markov_chain([0,0,0])
sage: m.next()
                        # random
{1: 0, 2: 0, 3: 0}
sage: m.next().values() # random
[0, 0, 0]
sage: m.next().values() # random
[0, 0, 0]
sage: m.next().values() # random
[0, 0, 0]
sage: m.next().values() # random
[0, 1, 0]
sage: m.next().values() # random
[0, 2, 0]
sage: m.next().values() # random
[0, 2, 1]
sage: m.next().values() # random
[1, 2, 1]
sage: m.next().values() # random
[2, 2, 1]
sage: m = s.markov_chain(s.zero_div(), [0.1,0.1,0.1,0.7])
sage: m.next().values() # random
[0, 0, 0, 1]
sage: m.next().values() # random
[0, 0, 1, 1]
sage: m.next().values() # random
[0, 0, 1, 2]
sage: m.next().values() # random
[1, 1, 2, 0]
sage: m.next().values() # random
[1, 1, 2, 1]
sage: m.next().values() # random
[1, 1, 2, 2]
```

```
sage: m.next().values() # random
[1, 1, 2, 3]
sage: m.next().values() # random
[1, 1, 2, 4]
sage: m.next().values() # random
[1, 1, 3, 4]
```

Note: The closed sandpile Markov chain has state space consisting of the configurations on a sandpile. It transitions from a state by choosing a vertex at random (according to the probability distribution distrib), dropping a grain of sand at that vertex, and stabilizing. If the chosen vertex is the sink, the chain stays at the current state.

The open sandpile Markov chain has state space consisting of the recurrent elements, i.e., the state space is the sandpile group. It transitions from the configuration c by choosing a vertex v at random according to distrib. The next state is the stabilization of c + v. If v is the sink vertex, then the stabilization of c + v is defined to be c.

Note that in either case, if distrib is specified, its length is equal to the total number of vertices (including the sink).

REFERENCES:

— max stable()

The maximal stable configuration.

OUTPUT:

SandpileConfig (the maximal stable configuration)

EXAMPLES:

```
sage: S = sandpiles.House()
sage: S.max_stable()
{1: 1, 2: 2, 3: 2, 4: 1}
```

— max_stable_div()

The maximal stable divisor.

OUTPUT:

SandpileDivisor (the maximal stable divisor)

EXAMPLES:

```
sage: s = sandpiles.Diamond()
sage: s.max_stable_div()
{0: 1, 1: 2, 2: 2, 3: 1}
sage: s.out_degree()
{0: 2, 1: 3, 2: 3, 3: 2}
```

— max_superstables(verbose=True)

The maximal superstable configurations. If the underlying graph is undirected, these are the superstables of highest degree. If verbose is False, the configurations are converted to lists of integers.

INPUT:

```
verbose - (default: True) boolean
```

OUTPUT:

tuple of SandpileConfig

```
sage: s = sandpiles.Diamond()
sage: s.superstables(False)
[[0, 0, 0],
  [0, 0, 1],
  [1, 0, 1],
  [0, 2, 0],
  [2, 0, 0],
  [0, 1, 1],
  [1, 0, 0],
  [0, 1, 0]]
sage: s.max_superstables(False)
[[1, 0, 1], [0, 2, 0], [2, 0, 0], [0, 1, 1]]
sage: s.h_vector()
[1, 3, 4]
```

— min_recurrents(verbose=True)

The minimal recurrent elements. If the underlying graph is undirected, these are the recurrent elements of least degree. If verbose is False, the configurations are converted to lists of integers.

INPUT:

```
verbose - (default: True) boolean
```

OUTPUT:

list of SandpileConfig

EXAMPLES:

```
sage: s = sandpiles.Diamond()
sage: s.recurrents(False)
[[2, 2, 1],
      [2, 2, 0],
      [1, 2, 0],
      [2, 0, 1],
      [0, 2, 1],
      [2, 1, 0],
      [1, 2, 1],
      [2, 1, 1]]
sage: s.min_recurrents(False)
[[1, 2, 0], [2, 0, 1], [0, 2, 1], [2, 1, 0]]
sage: [i.deg() for i in s.recurrents()]
[5, 4, 3, 3, 3, 3, 4, 4]
```

— nonsink vertices()

The nonsink vertices.

OUTPUT:

list of vertices

EXAMPLES:

```
sage: s = sandpiles.Grid(2,3)
sage: s.nonsink_vertices()
[(1, 1), (1, 2), (1, 3), (2, 1), (2, 2), (2, 3)]
```

— nonspecial_divisors(verbose=True)

The nonspecial divisors. Only for undirected graphs. (See NOTE.)

INPUT:

verbose - (default: True) boolean

OUTPUT:

list (of divisors)

EXAMPLES:

```
sage: S = sandpiles.Complete(4)
sage: ns = S.nonspecial_divisors()
sage: D = ns[0]
sage: D.values()
[-1, 0, 1, 2]
sage: D.deg()
2
sage: [i.effective_div() for i in ns]
[[], [], [], [], [], []]
```

Note: The "nonspecial divisors" are those divisors of degree g-1 with empty linear system. The term is only defined for undirected graphs. Here, g=|E|-|V|+1 is the genus of the graph (not counted loops as part of |E|). If verbose is False, the divisors are converted to lists of integers.

Warning: The underlying graph must be undirected.

— out_degree(v=None)

The out-degree of a vertex or a list of all out-degrees.

INPUT:

v - (optional) vertex name

OUTPUT:

integer or dict

EXAMPLES:

```
sage: s = sandpiles.House()
sage: s.out_degree()
{0: 2, 1: 2, 2: 3, 3: 3, 4: 2}
sage: s.out_degree(2)
```

— picard_representatives(d, verbose=True)

Representatives of the divisor classes of degree d in the Picard group. (Also see the documentation for jacobian_representatives.)

INPUT:

- d integer
- verbose (default: True) boolean

OUTPUT:

list of SandpileDivisors (or lists representing divisors)

```
sage: s = sandpiles.Complete(3)
sage: s.superstables(False)
[[0, 0], [0, 1], [1, 0]]
sage: s.jacobian_representatives(False)
[[0, 0, 0], [-1, 0, 1], [-1, 1, 0]]
sage: s.picard_representatives(3,False)
[[3, 0, 0], [2, 0, 1], [2, 1, 0]]
```

— points()

Generators for the multiplicative group of zeros of the sandpile ideal.

OUTPUT:

list of complex numbers

EXAMPLES:

The sandpile group in this example is cyclic, and hence there is a single generator for the group of solutions.

```
sage: S = sandpiles.Complete(4)
sage: S.points()
[[1, I, -I], [I, 1, -I]]
```

— postulation()

The postulation number of the toppling ideal. This is the largest weight of a superstable configuration of the graph.

OUTPUT:

nonnegative integer

EXAMPLES:

```
sage: s = sandpiles.Complete(4)
sage: s.postulation()
3
```

— recurrents(verbose=True)

The recurrent configurations. If verbose is False, the configurations are converted to lists of integers.

INPUT:

```
verbose - (default: True) boolean
```

OUTPUT:

list of recurrent configurations

```
[1, 2, 1],
[1, 1, 2],
[2, 1, 0],
[2, 0, 1],
[1, 2, 0],
[1, 0, 2],
[0, 2, 1],
[0, 1, 2]]
sage: sandpiles.Cycle(4).recurrents(False)
[[1, 1, 1], [0, 1, 1], [1, 0, 1], [1, 1, 0]]
```

— reduced_laplacian()

The reduced Laplacian matrix of the graph.

OUTPUT:

matrix

EXAMPLES:

```
sage: S = sandpiles.Diamond()
sage: S.laplacian()
[ 2 -1 -1 0]
[-1 3 -1 -1]
[-1 -1 3 -1]
[ 0 -1 -1 2]
sage: S.reduced_laplacian()
[ 3 -1 -1]
[-1 3 -1]
[-1 -1 2]
```

Note: This is the Laplacian matrix with the row and column indexed by the sink vertex removed.

— reorder_vertices()

A copy of the sandpile with vertex names permuted. After reordering, vertex u comes before vertex v in the list of vertices if u is closer to the sink.

OUTPUT:

Sandpile

EXAMPLES:

```
sage: S = Sandpile({0:[1], 2:[0,1], 1:[2]})
sage: S.dict()
{0: {1: 1}, 1: {2: 1}, 2: {0: 1, 1: 1}}
sage: T = S.reorder_vertices()
```

The vertices 1 and 2 have been swapped:

```
sage: T.dict()
{0: {1: 1}, 1: {0: 1, 2: 1}, 2: {0: 1}}
```

— resolution(verbose=False)

A minimal free resolution of the homogeneous toppling ideal. If verbose is True, then all of the mappings are returned. Otherwise, the resolution is summarized.

INPUT:

verbose - (default: False) boolean

OUTPUT:

free resolution of the toppling ideal

EXAMPLES:

```
sage: S = Sandpile({0: {}}, 1: {0: 1, 2: 1, 3: 4}, 2: {3: 5}, 3: {1: 1, 2: 1}},0)
sage: S.resolution() # a Gorenstein sandpile graph
'R^1 <-- R^5 <-- R^5 <-- R^1'
sage: S.resolution(True)
[x1^2 - x3*x0 \ x3*x1 - x2*x0 \ x3^2 - x2*x1 \ x2*x3 - x0^2 \ x2^2 - x1*x0],
[ x3 x2 0 x0 0] [ x2^2 - x1*x0]
[-x1 -x3  x2  0 -x0]  [-x2*x3 + x0^2]
[ x0 x1 0 x2 0] [-x3^2 + x2*x1]
[ 0 0 -x1 -x3 x2] [x3*x1 - x2*x0]
\begin{bmatrix} 0 & 0 & x0 & x1 - x3 \end{bmatrix}, \begin{bmatrix} x1^2 - x3 \times x0 \end{bmatrix}
sage: r = S.resolution(True)
sage: r[0]*r[1]
[0 0 0 0 0]
sage: r[1]*r[2]
[0]
[0]
[0]
[0]
[0]
```

- ring()

The ring containing the homogeneous toppling ideal.

OUTPUT:

ring

EXAMPLES:

```
sage: S = sandpiles.Diamond()
sage: S.ring()
Multivariate Polynomial Ring in x3, x2, x1, x0 over Rational Field
sage: S.ring().gens()
(x3, x2, x1, x0)
```

Note: The indeterminate xi corresponds to the i-th vertex as listed my the method vertices. The term-ordering is degrevlex with indeterminates ordered according to their distance from the sink (larger indeterminates are further from the sink).

- show(**kwds)

Draw the underlying graph.

INPUT:

kwds – (optional) arguments passed to the show method for Graph or DiGraph

```
sage: S = Sandpile({0:[], 1:[0,3,4], 2:[0,3,5], 3:[2,5], 4:[1,1], 5:[2,4]})
sage: S.show()
sage: S.show(graph_border=True, edge_labels=True)
```

— show3d(**kwds)

Draw the underlying graph.

INPUT:

kwds – (optional) arguments passed to the show method for Graph or DiGraph

EXAMPLES:

```
sage: S = sandpiles.House()
sage: S.show3d()
```

— sink()

The sink vertex.

OUTPUT:

sink vertex

EXAMPLES:

```
sage: G = sandpiles.House()
sage: G.sink()
0
sage: H = sandpiles.Grid(2,2)
sage: H.sink()
(0, 0)
sage: type(H.sink())
<type 'tuple'>
```

- smith_form()

The Smith normal form for the Laplacian. In detail: a list of integer matrices D, U, V such that ULV = D where L is the transpose of the Laplacian, D is diagonal, and U and V are invertible over the integers.

OUTPUT:

list of integer matrices

EXAMPLES:

```
sage: s = sandpiles.Complete(4)
sage: D,U,V = s.smith_form()
sage: D
[1 0 0 0]
[0 4 0 0]
[0 0 4 0]
[0 0 0 0]
sage: U*s.laplacian()*V == D # laplacian symmetric => tranpose not necessary
True
```

— solve()

Approximations of the complex affine zeros of the sandpile ideal.

OUTPUT:

list of complex numbers

EXAMPLES:

```
sage: S = Sandpile({0: {}, 1: {2: 2}, 2: {0: 4, 1: 1}}, 0)
sage: S.solve()
[[-0.707107 + 0.707107*I, 0.707107 - 0.707107*I], [-0.707107 - 0.707107*I, 0.707107 + 0.707107*I], [-sage: len(_)
8
sage: S.group_order()
8
```

Note: The solutions form a multiplicative group isomorphic to the sandpile group. Generators for this group are given exactly by points().

— stable_configs(smax=None)

Generator for all stable configurations. If smax is provided, then the generator gives all stable configurations less than or equal to smax. If smax does not represent a stable configuration, then each component of smax is replaced by the corresponding component of the maximal stable configuration.

INPUT:

smax – (optional) SandpileConfig or list representing a SandpileConfig

OUTPUT:

generator for all stable configurations

EXAMPLES:

```
sage: s = sandpiles.Complete(3)
sage: a = s.stable_configs()
sage: a.next()
{1: 0, 2: 0}
sage: [i.values() for i in a]
[[0, 1], [1, 0], [1, 1]]
sage: b = s.stable_configs([1,0])
sage: list(b)
[{1: 0, 2: 0}, {1: 1, 2: 0}]
```

— stationary_density()

The stationary density of the sandpile.

OUTPUT:

rational number

EXAMPLES:

```
sage: s = sandpiles.Complete(3)
sage: s.stationary_density()
10/9
sage: s = Sandpile(digraphs.DeBruijn(2,2),'00')
sage: s.stationary_density()
9/8
```

Note: The stationary density of a sandpile is the sum $\sum_c (\deg(c) + \deg(s))$ where $\deg(s)$ is the degree of the sink and the sum is over all recurrent configurations.

REFERENCES:

— superstables(verbose=True)

The superstable configurations. If verbose is False, the configurations are converted to lists of integers. Superstables for undirected graphs are also known as G-parking functions.

INPUT:

```
verbose - (default: True) boolean
```

OUTPUT:

list of SandpileConfig

EXAMPLES:

```
sage: sp = Sandpile(graphs.HouseXGraph(),0).superstables()
sage: sp[:3]
[\{1: 0, 2: 0, 3: 0, 4: 0\}, \{1: 1, 2: 0, 3: 0, 4: 1\}, \{1: 1, 2: 0, 3: 0, 4: 0\}]
sage: sandpiles.Complete(4).superstables(False)
[[0, 0, 0],
[0, 0, 1],
 [0, 1, 0],
 [1, 0, 0],
 [0, 0, 2],
 [0, 2, 0],
 [2, 0, 0],
 [0, 1, 1],
 [1, 0, 1],
 [1, 1, 0],
 [0, 1, 2],
 [0, 2, 1],
 [1, 0, 2],
 [1, 2, 0],
 [2, 0, 1],
 [2, 1, 0]]
sage: sandpiles.Cycle(4).superstables(False)
[[0, 0, 0], [1, 0, 0], [0, 1, 0], [0, 0, 1]]
```

— symmetric_recurrents(orbits)

The symmetric recurrent configurations.

INPUT:

orbits - list of lists partitioning the vertices

OUTPUT:

list of recurrent configurations

```
{1: 2, 2: 2, 3: 2, 4: 0},

{1: 2, 2: 1, 3: 2, 4: 0},

{1: 2, 2: 2, 3: 0, 4: 1},

{1: 2, 2: 0, 3: 2, 4: 1},

{1: 2, 2: 2, 3: 1, 4: 0},

{1: 2, 2: 1, 3: 2, 4: 1},

{1: 2, 2: 2, 3: 1, 4: 1}
```

Note: The user is responsible for ensuring that the list of orbits comes from a group of symmetries of the underlying graph.

— tutte_polynomial()

The Tutte polynomial. Only defined for undirected sandpile graphs.

OUTPUT:

polynomial

EXAMPLES:

```
sage: s = sandpiles.Complete(4)
sage: s.tutte_polynomial()
x^3 + y^3 + 3*x^2 + 4*x*y + 3*y^2 + 2*x + 2*y
sage: s.tutte_polynomial().subs(x=1)
y^3 + 3*y^2 + 6*y + 6
sage: s.tutte_polynomial().subs(x=1).coefficients() == s.h_vector()
True
```

— unsaturated_ideal()

The unsaturated, homogeneous toppling ideal.

OUTPUT:

ideal

EXAMPLES:

```
sage: S = sandpiles.Diamond()
sage: S.unsaturated_ideal().gens()
[x1^3 - x3*x2*x0, x2^3 - x3*x1*x0, x3^2 - x2*x1]
sage: S.ideal().gens()
[x2*x1 - x0^2, x3^2 - x0^2, x1^3 - x3*x2*x0, x3*x1^2 - x2^2*x0, x2^3 - x3*x1*x0, x3*x2^2 - x1^2*x0]
```

— version()

The version number of Sage Sandpiles.

OUTPUT:

string

```
sage: Sandpile.version()
Sage Sandpiles Version 2.4
sage: S = sandpiles.Complete(3)
sage: S.version()
Sage Sandpiles Version 2.4
```

— zero_config()

The all-zero configuration.

OUTPUT:

SandpileConfig

EXAMPLES:

```
sage: s = sandpiles.Diamond()
sage: s.zero_config()
{1: 0, 2: 0, 3: 0}
```

— zero_div()

The all-zero divisor.

OUTPUT:

SandpileDivisor

EXAMPLES:

```
sage: S = sandpiles.House()
sage: S.zero_div()
{0: 0, 1: 0, 2: 0, 3: 0, 4: 0}
```

SandpileConfig Summary of methods.

- + Addition of configurations.
- & The stabilization of the sum.
- greater-equal True if every component of self is at least that of other.
- greater True if every component of self is at least that of other and the two configurations are not equal.
- ~ The stabilized configuration.
- less-equal True if every component of self is at most that of other.
- less True if every component of self is at most that of other and the two configurations are not equal.
- * The recurrent element equivalent to the sum.
- ^ Exponentiation for the *-operator.
- - The additive inverse of the configuration.
- - Subtraction of configurations.
- add_random Add one grain of sand to a random vertex.
- burst_size The burst size of the configuration with respect to the given vertex.
- deg The degree of the configuration.
- dualize The difference with the maximal stable configuration.
- equivalent_recurrent The recurrent configuration equivalent to the given configuration.
- equivalent_superstable The equivalent superstable configuration.
- fire_script Fire the given script.

- fire_unstable Fire all unstable vertices.
- fire_vertex Fire the given vertex.
- *help* List of SandpileConfig methods.
- *is_recurrent* Is the configuration recurrent?
- is stable Is the configuration stable?
- *is_superstable* Is the configuration superstable?
- *is_symmetric* Is the configuration symmetric?
- order The order of the equivalent recurrent element.
- *sandpile* The configuration's underlying sandpile.
- *show* Show the configuration.
- *stabilize* The stabilized configuration.
- support The vertices containing sand.
- unstable The unstable vertices.
- values The values of the configuration as a list.

Complete descriptions of SandpileConfig methods.

```
Addition of configurations.
    INPUT:
    other - SandpileConfig
    OUTPUT:
    sum of self and other
    EXAMPLES:
    sage: S = sandpiles.Cycle(3)
    sage: c = SandpileConfig(S, [1,2])
    sage: d = SandpileConfig(S, [3,2])
    sage: c + d
     {1: 4, 2: 4}
— &
    The stabilization of the sum.
    INPUT:
    other - SandpileConfig
    OUTPUT:
    SandpileConfig
    EXAMPLES:
```

```
sage: S = sandpiles.Cycle(4)
    sage: c = SandpileConfig(S, [1,0,0])
    sage: c + c # ordinary addition
     {1: 2, 2: 0, 3: 0}
    sage: c & c # add and stabilize
     {1: 0, 2: 1, 3: 0}
    sage: c*c # add and find equivalent recurrent
    {1: 1, 2: 1, 3: 1}
    sage: ~(c + c) == c & c
    True
-- >=
    True if every component of self is at least that of other.
    INPUT:
    other - SandpileConfig
    OUTPUT:
    boolean
    EXAMPLES:
    sage: S = sandpiles.Cycle(3)
    sage: c = SandpileConfig(S, [1,2])
    sage: d = SandpileConfig(S, [2,3])
    sage: e = SandpileConfig(S, [2,0])
    sage: c >= c
    True
    sage: d >= c
    True
    sage: c >= d
    False
    sage: e >= c
    False
    sage: c >= e
    False
-- >
    True if every component of self is at least that of other and the two configurations are not equal.
    INPUT:
    other - SandpileConfig
    OUTPUT:
    boolean
    EXAMPLES:
    sage: S = sandpiles.Cycle(3)
    sage: c = SandpileConfig(S, [1,2])
    sage: d = SandpileConfig(S, [1,3])
    sage: c > c
    False
    sage: d > c
    True
```

sage: c > d
False

```
The stabilized configuration.
     OUTPUT:
     SandpileConfig
     EXAMPLES:
     sage: S = sandpiles.House()
     sage: c = S.max_stable() + S.identity()
     sage: ~c == c.stabilize()
     True
-- <=
     True if every component of self is at most that of other.
     INPUT:
     other - SandpileConfig
     OUTPUT:
     boolean
     EXAMPLES:
     sage: S = sandpiles.Cycle(3)
     sage: c = SandpileConfig(S, [1,2])
     sage: d = SandpileConfig(S, [2,3])
     sage: e = SandpileConfig(S, [2,0])
     sage: c <= c</pre>
     True
     sage: c <= d</pre>
     True
     sage: d <= c</pre>
     False
     sage: c <= e</pre>
     False
     sage: e <= c</pre>
     False
— <
     True if every component of self is at most that of other and the two configurations are not equal.
     INPUT:
     other-SandpileConfig
     OUTPUT:
     boolean
     EXAMPLES:
     sage: S = sandpiles.Cycle(3)
     sage: c = SandpileConfig(S, [1,2])
     sage: d = SandpileConfig(S, [2,3])
     sage: c < c</pre>
     False
```

sage: c < d</pre>

True

```
sage: d < c
False
sage: S = Sandpile(graphs.CycleGraph(3), 0)
sage: c = SandpileConfig(S, [1,2])
sage: d = SandpileConfig(S, [2,3])
sage: c < c
False
sage: c < d
True
sage: d < c
False</pre>
```

If other is an configuration, the recurrent element equivalent to the sum. If other is an integer, the sum of configuration with itself other times.

INPUT:

other - SandpileConfig or Integer

OUTPUT:

SandpileConfig

EXAMPLES:

```
sage: S = sandpiles.Cycle(4)
sage: c = SandpileConfig(S, [1,0,0])
sage: c + c # ordinary addition
{1: 2, 2: 0, 3: 0}
sage: c & c # add and stabilize
{1: 0, 2: 1, 3: 0}
sage: c*c # add and find equivalent recurrent
{1: 1, 2: 1, 3: 1}
sage: (c*c).is_recurrent()
True
sage: c*(-c) == S.identity()
True
sage: c
{1: 1, 2: 0, 3: 0}
sage: c*3
{1: 3, 2: 0, 3: 0}
```

The recurrent element equivalent to the sum of the configuration with itself k times. If k is negative, do the same for the negation of the configuration. If k is zero, return the identity of the sandpile group.

INPUT:

__ ^

k - SandpileConfig

OUTPUT:

SandpileConfig

```
sage: S = sandpiles.Cycle(4)
sage: c = SandpileConfig(S, [1,0,0])
sage: c^3
{1: 1, 2: 1, 3: 0}
```

```
sage: (c + c + c) == c^3
False
sage: (c + c + c).equivalent_recurrent() == c^3
sage: c^(-1)
{1: 1, 2: 1, 3: 0}
sage: c^0 == S.identity()
True
The additive inverse of the configuration.
OUTPUT:
SandpileConfig
EXAMPLES:
sage: S = sandpiles.Cycle(3)
sage: c = SandpileConfig(S, [1,2])
sage: -c
\{1: -1, 2: -2\}
Subtraction of configurations.
INPUT:
other-SandpileConfig
OUTPUT:
sum of self and other
EXAMPLES:
sage: S = sandpiles.Cycle(3)
```

— add_random(distrib=None)

sage: c - d
{1: -2, 2: 0}

sage: c = SandpileConfig(S, [1,2])
sage: d = SandpileConfig(S, [3,2])

Add one grain of sand to a random vertex. Optionally, a probability distribution, distrib, may be placed on the vertices or the nonsink vertices. See NOTE for details.

INPUT:

distrib – (optional) list of nonnegative numbers summing to 1 (representing a prob. dist.)

OUTPUT:

SandpileConfig

```
sage: s = sandpiles.Complete(4)
sage: c = s.zero_config()
sage: c.add_random() # random
{1: 0, 2: 1, 3: 0}
sage: c
{1: 0, 2: 0, 3: 0}
```

```
sage: c.add_random([0.1,0.1,0.8]) # random
{1: 0, 2: 0, 3: 1}
sage: c.add_random([0.7,0.1,0.1,0.1]) # random
{1: 0, 2: 0, 3: 0}
```

We compute the "sizes" of the avalanches caused by adding random grains of sand to the maximal stable configuration on a grid graph. The function stabilize() returns the firing vector of the stabilization, a dictionary whose values say how many times each vertex fires in the stabilization.:

Note: If distrib is None, then the probability is the uniform probability on the nonsink vertices. Otherwise, there are two possibilities:

- (i) the length of distrib is equal to the number of vertices, and distrib represents a probability distribution on all of the vertices. In that case, the sink may be chosen at random, in which case, the configuration is unchanged.
- (ii) Otherwise, the length of distrib must be equal to the number of nonsink vertices, and distrib represents a probability distribution on the nonsink vertices.

Warning: If distrib != None, the user is responsible for assuring the sum of its entries is 1 and that its length is equal to the number of sink vertices or the number of nonsink vertices.

— burst size(v)

The burst size of the configuration with respect to the given vertex.

INPUT:

v – vertex

OUTPUT:

integer

EXAMPLES:

```
sage: s = sandpiles.Diamond()
sage: [i.burst_size(0) for i in s.recurrents()]
[1, 1, 1, 1, 1, 1, 1]
sage: [i.burst_size(1) for i in s.recurrents()]
[0, 0, 1, 2, 1, 2, 0, 2]
```

Note: To define c.burst (v), if v is not the sink, let c' be the unique recurrent for which the stabilization of c' + v is c. The burst size is then the amount of sand that goes into the sink during this stabilization. If v is the sink, the burst size is defined to be 1.

REFERENCES:

— deg()

The degree of the configuration.

OUTPUT:

integer

EXAMPLES:

```
sage: S = sandpiles.Complete(3)
sage: c = SandpileConfig(S, [1,2])
sage: c.deg()
3
```

— dualize()

The difference with the maximal stable configuration.

OUTPUT:

SandpileConfig

EXAMPLES:

```
sage: S = sandpiles.Cycle(3)
sage: c = SandpileConfig(S, [1,2])
sage: S.max_stable()
{1: 1, 2: 1}
sage: c.dualize()
{1: 0, 2: -1}
sage: S.max_stable() - c == c.dualize()
```

— equivalent_recurrent(with_firing_vector=False)

The recurrent configuration equivalent to the given configuration. Optionally, return the corresponding firing vector.

INPUT:

```
with_firing_vector - (default: False) boolean
```

OUTPUT:

SandpileConfig or [SandpileConfig, firing_vector]

EXAMPLES:

```
sage: S = sandpiles.Diamond()
sage: c = SandpileConfig(S, [0,0,0])
sage: c.equivalent_recurrent() == S.identity()
True
sage: x = c.equivalent_recurrent(True)
sage: r = vector([x[0][v] for v in S.nonsink_vertices()])
sage: f = vector([x[1][v] for v in S.nonsink_vertices()])
sage: cv = vector(c.values())
sage: r == cv - f*S.reduced_laplacian()
```

Note: Let L be the reduced Laplacian, c the initial configuration, r the returned configuration, and f the firing vector. Then $r = c - f \cdot L$.

— equivalent_superstable(with_firing_vector=False)

The equivalent superstable configuration. Optionally, return the corresponding firing vector.

INPUT:

```
with_firing_vector - (default: False) boolean
```

OUTPUT:

SandpileConfig or [SandpileConfig, firing vector]

EXAMPLES:

```
sage: S = sandpiles.Diamond()
sage: m = S.max_stable()
sage: m.equivalent_superstable().is_superstable()
True
sage: x = m.equivalent_superstable(True)
sage: s = vector(x[0].values())
sage: f = vector(x[1].values())
sage: mv = vector(m.values())
sage: s == mv - f*S.reduced_laplacian()
True
```

Note: Let L be the reduced Laplacian, c the initial configuration, s the returned configuration, and f the firing vector. Then $s = c - f \cdot L$.

— fire script(sigma)

Fire the given script. In other words, fire each vertex the number of times indicated by sigma.

INPUT:

sigma - SandpileConfig or (list or dict representing a SandpileConfig)

OUTPUT:

SandpileConfig

EXAMPLES:

```
sage: S = sandpiles.Cycle(4)
sage: c = SandpileConfig(S, [1,2,3])
sage: c.unstable()
[2, 3]
sage: c.fire_script(SandpileConfig(S,[0,1,1]))
{1: 2, 2: 1, 3: 2}
sage: c.fire_script(SandpileConfig(S,[2,0,0])) == c.fire_vertex(1).fire_vertex(1)
True
```

— fire_unstable()

Fire all unstable vertices.

OUTPUT:

SandpileConfig

```
sage: S = sandpiles.Cycle(4)
sage: c = SandpileConfig(S, [1,2,3])
```

```
sage: c.fire_unstable()
{1: 2, 2: 1, 3: 2}
— fire vertex(v)
Fire the given vertex.
INPUT:
v - vertex
OUTPUT:
SandpileConfig
EXAMPLES:
sage: S = sandpiles.Cycle(3)
sage: c = SandpileConfig(S, [1,2])
sage: c.fire_vertex(2)
{1: 2, 2: 0}
— help(verbose=True)
List of SandpileConfig methods. If verbose, include short descriptions.
INPUT:
verbose - (default: True) boolean
OUTPUT:
printed string
EXAMPLES:
sage: SandpileConfig.help()
Shortcuts for SandpileConfig operations:
~c -- stabilize
c & d -- add and stabilize
c \, \star \, c \, -- \, add and find equivalent recurrent
c^k -- add k times and find equivalent recurrent
         (taking inverse if k is negative)
For detailed help with any method FOO listed below,
enter "SandpileConfig.FOO?" or enter "c.FOO?" for any SandpileConfig c.
                       -- Add one grain of sand to a random vertex.
add_random
burst_size
                       -- The burst size of the configuration with respect to the given vertex.
deg
                       -- The degree of the configuration.
                      -- The difference with the maximal stable configuration.
equivalent_recurrent -- The recurrent configuration equivalent to the given configuration.
equivalent_superstable -- The equivalent superstable configuration.
fire_script -- Fire the given script.
                      -- Fire all unstable vertices.
fire_unstable
fire_vertex
                     -- Fire the given vertex.
                       -- List of SandpileConfig methods.
help
is_recurrent -- Is the configuration recurrent?
is_stable
                       -- Is the configuration stable?
is_superstable
is_symmetric
                     -- Is the configuration superstable?
                      -- Is the configuration symmetric?
order
                       -- The order of the equivalent recurrent element.
```

```
sandpile -- The configuration's underlying sandpile.
show -- Show the configuration.
stabilize -- The stabilized configuration.
support -- The vertices containing sand.
unstable -- The unstable vertices.
values -- The values of the configuration as a list.
```

— is_recurrent()

Is the configuration recurrent?

OUTPUT:

boolean

EXAMPLES:

```
sage: S = sandpiles.Diamond()
sage: S.identity().is_recurrent()
True
sage: S.zero_config().is_recurrent()
False
```

— is stable()

Is the configuration stable?

OUTPUT:

boolean

EXAMPLES:

```
sage: S = sandpiles.Diamond()
sage: S.max_stable().is_stable()
True
sage: (2*S.max_stable()).is_stable()
False
sage: (S.max_stable() & S.max_stable()).is_stable()
True
```

— is_superstable()

Is the configuration superstable?

OUTPUT:

boolean

EXAMPLES:

```
sage: S = sandpiles.Diamond()
sage: S.zero_config().is_superstable()
True
```

— is_symmetric(orbits)

Is the configuration symmetric? Return True if the values of the configuration are constant over the vertices in each sublist of orbits.

INPUT:

orbits - list of lists of vertices

OUTPUT:

boolean

EXAMPLES:

— order()

The order of the equivalent recurrent element.

OUTPUT:

integer

EXAMPLES:

```
sage: S = sandpiles.Diamond()
sage: c = SandpileConfig(S,[2,0,1])
sage: c.order()
4
sage: ~(c + c + c + c) == S.identity()
True
sage: c = SandpileConfig(S,[1,1,0])
sage: c.order()
1
sage: c.is_recurrent()
False
sage: c.equivalent_recurrent() == S.identity()
True
```

— sandpile()

The configuration's underlying sandpile.

OUTPUT:

Sandpile

EXAMPLES:

```
sage: S = sandpiles.Diamond()
sage: c = S.identity()
sage: c.sandpile()
Diamond sandpile graph: 4 vertices, sink = 0
sage: c.sandpile() == S
True
```

— show(sink=True, colors=True, heights=False, directed=None, **kwds)

Show the configuration.

INPUT:

• sink – (default: True) whether to show the sink

- colors (default: True) whether to color-code the amount of sand on each vertex
- heights (default: False) whether to label each vertex with the amount of sand
- directed (optional) whether to draw directed edges
- kwds (optional) arguments passed to the show method for Graph

EXAMPLES:

```
sage: S = sandpiles.Diamond()
sage: c = S.identity()
sage: c.show()
sage: c.show(directed=False)
sage: c.show(sink=False,colors=False,heights=True)
```

— stabilize(with_firing_vector=False)

The stabilized configuration. Optionally returns the corresponding firing vector.

INPUT:

```
with_firing_vector - (default: False) boolean
```

OUTPUT:

SandpileConfig or [SandpileConfig, firing_vector]

EXAMPLES:

```
sage: S = sandpiles.House()
sage: c = 2*S.max_stable()
sage: c._set_stabilize()
sage: '_stabilize' in c.__dict__
True
sage: S = sandpiles.House()
sage: c = S.max_stable() + S.identity()
sage: c.stabilize(True)
[{1: 1, 2: 2, 3: 2, 4: 1}, {1: 2, 2: 2, 3: 3, 4: 3}]
sage: S.max_stable() & S.identity() == c.stabilize()
True
sage: ~c == c.stabilize()
```

— support()

The vertices containing sand.

OUTPUT:

list - support of the configuration

EXAMPLES:

```
sage: S = sandpiles.Diamond()
sage: c = S.identity()
sage: c
{1: 2, 2: 2, 3: 0}
sage: c.support()
[1, 2]
```

— unstable()

The unstable vertices.

OUTPUT:

list of vertices

EXAMPLES:

```
sage: S = sandpiles.Cycle(4)
sage: c = SandpileConfig(S, [1,2,3])
sage: c.unstable()
[2, 3]
```

— values()

The values of the configuration as a list. The list is sorted in the order of the vertices.

OUTPUT:

list of integers

boolean

EXAMPLES:

```
sage: S = Sandpile({'a':[1,'b'], 'b':[1,'a'], 1:['a']},'a')
sage: c = SandpileConfig(S, {'b':1, 1:2})
sage: c
{1: 2, 'b': 1}
sage: c.values()
[2, 1]
sage: S.nonsink_vertices()
[1, 'b']
```

SandpileDivisor Summary of methods.

- + Addition of divisors.
- greater-equal True if every component of self is at least that of other.
- greater True if every component of self is at least that of other and the two divisors are not equal.
- less-equal True if every component of self is at most that of other.
- less True if every component of self is at most that of other and the two divisors are not equal.
- - The additive inverse of the divisor.
- - Subtraction of divisors.
- *Dcomplex* The support-complex.
- add_random Add one grain of sand to a random vertex.
- betti The Betti numbers for the support-complex.
- deg The degree of the divisor.
- dualize The difference with the maximal stable divisor.
- effective_div All linearly equivalent effective divisors.
- fire_script Fire the given script.
- fire_unstable Fire all unstable vertices.

- fire_vertex Fire the given vertex.
- *help* List of SandpileDivisor methods.
- *is_alive* Is the divisor stabilizable?
- is_linearly_equivalent Is the given divisor linearly equivalent?
- is q reduced Is the divisor q-reduced?
- is_symmetric Is the divisor symmetric?
- is_weierstrass_pt Is the given vertex a Weierstrass point?
- *polytope* The polytope determining the complete linear system.
- polytope_integer_pts The integer points inside divisor's polytope.
- *q_reduced* The linearly equivalent q-reduced divisor.
- rank The rank of the divisor.
- sandpile The divisor's underlying sandpile.
- *show* Show the divisor.
- simulate threshold The first unstabilizable divisor in the closed Markov chain.
- stabilize The stabilization of the divisor.
- *support* List of vertices at which the divisor is nonzero.
- unstable The unstable vertices.
- values The values of the divisor as a list.
- weierstrass_div The Weierstrass divisor.
- weierstrass_gap_seq The Weierstrass gap sequence at the given vertex.
- weierstrass_pts The Weierstrass points (vertices).
- weierstrass_rank_seq The Weierstrass rank sequence at the given vertex.

Complete descriptions of SandpileDivisor methods.

```
Addition of divisors.

INPUT:

other - SandpileDivisor

OUTPUT:

sum of self and other

EXAMPLES:

sage: S = sandpiles.Cycle(3)

sage: D = SandpileDivisor(S, [1,2,3])

sage: E = SandpileDivisor(S, [3,2,1])

sage: D + E

{0: 4, 1: 4, 2: 4}
```

-- >=

```
True if every component of self is at least that of other.
     INPUT:
     other - SandpileDivisor
     OUTPUT:
     boolean
     EXAMPLES:
     sage: S = sandpiles.Cycle(3)
     sage: D = SandpileDivisor(S, [1,2,3])
     sage: E = SandpileDivisor(S, [2,3,4])
     sage: F = SandpileDivisor(S, [2,0,4])
     sage: D >= D
     True
     sage: E >= D
     True
     sage: D >= E
     False
     sage: F >= D
    False
     sage: D >= F
    False
-- >
     True if every component of self is at least that of other and the two divisors are not equal.
     INPUT:
     other - SandpileDivisor
     OUTPUT:
     boolean
     EXAMPLES:
     sage: S = sandpiles.Cycle(3)
     sage: D = SandpileDivisor(S, [1,2,3])
     sage: E = SandpileDivisor(S, [1,3,4])
     sage: D > D
     False
     sage: E > D
    True
     sage: D > E
    False
-- <=
     True if every component of self is at most that of other.
    INPUT:
     other - SandpileDivisor
     OUTPUT:
     boolean
```

```
sage: S = sandpiles.Cycle(3)
sage: D = SandpileDivisor(S, [1,2,3])
sage: E = SandpileDivisor(S, [2,3,4])
sage: F = SandpileDivisor(S, [2, 0, 4])
sage: D <= D</pre>
True
sage: D <= E</pre>
True
sage: E <= D</pre>
False
sage: D <= F</pre>
False
sage: F <= D</pre>
False
True if every component of self is at most that of other and the two divisors are not equal.
INPUT:
other - SandpileDivisor
OUTPUT:
boolean
EXAMPLES:
sage: S = sandpiles.Cycle(3)
sage: D = SandpileDivisor(S, [1,2,3])
sage: E = SandpileDivisor(S, [2,3,4])
sage: D < D</pre>
False
sage: D < E</pre>
True
sage: E < D</pre>
False
The additive inverse of the divisor.
OUTPUT:
SandpileDivisor
EXAMPLES:
sage: S = sandpiles.Cycle(3)
sage: D = SandpileDivisor(S, [1,2,3])
sage: -D
\{0: -1, 1: -2, 2: -3\}
Subtraction of divisors.
INPUT:
other - SandpileDivisor
OUTPUT:
Difference of self and other
```

EXAMPLES:

```
sage: S = sandpiles.Cycle(3)
sage: D = SandpileDivisor(S, [1,2,3])
sage: E = SandpileDivisor(S, [3,2,1])
sage: D - E
{0: -2, 1: 0, 2: 2}
```

— Dcomplex()

The support-complex. (See NOTE.)

OUTPUT:

simplicial complex

EXAMPLES:

```
sage: S = sandpiles.House()
sage: p = SandpileDivisor(S, [1,2,1,0,0]).Dcomplex()
sage: p.homology()
{0: 0, 1: Z x Z, 2: 0}
sage: p.f_vector()
[1, 5, 10, 4]
sage: p.betti()
{0: 1, 1: 2, 2: 0}
```

Note: The "support-complex" is the simplicial complex determined by the supports of the linearly equivalent effective divisors.

— add_random(distrib=None)

Add one grain of sand to a random vertex.

INPUT:

distrib - (optional) list of nonnegative numbers representing a probability distribution on the vertices

OUTPUT:

SandpileDivisor

EXAMPLES:

```
sage: s = sandpiles.Complete(4)
sage: D = s.zero_div()
sage: D.add_random() # random
{0: 0, 1: 0, 2: 1, 3: 0}
sage: D.add_random([0.1,0.1,0.1,0.7]) # random
{0: 0, 1: 0, 2: 0, 3: 1}
```

Warning: If distrib is not None, the user is responsible for assuring the sum of its entries is 1.

- betti()

The Betti numbers for the support-complex. (See NOTE.)

OUTPUT:

dictionary of integers

```
sage: S = sandpiles.Cycle(3)
sage: D = SandpileDivisor(S, [2,0,1])
sage: D.betti()
{0: 1, 1: 1}
```

Note: The "support-complex" is the simplicial complex determined by the supports of the linearly equivalent effective divisors.

— deg()

The degree of the divisor.

OUTPUT:

integer

EXAMPLES:

```
sage: S = sandpiles.Cycle(3)
sage: D = SandpileDivisor(S, [1,2,3])
sage: D.deg()
6
```

— dualize()

The difference with the maximal stable divisor.

OUTPUT:

SandpileDivisor

```
EXAMPLES:: sage: S = sandpiles.Cycle(3) sage: D = SandpileDivisor(S, [1,2,3]) sage: D.dualize() {0: 0, 1: -1, 2: -2} sage: S.max_stable_div() - D == D.dualize() True
```

$-- effective_div(verbose=True, with_firing_vectors=False)\\$

All linearly equivalent effective divisors. If verbose is False, the divisors are converted to lists of integers. If with_firing_vectors is True then a list of firing vectors is also given, each of which prescribes the vertices to be fired in order to obtain an effective divisor.

INPUT:

- verbose (default: True) boolean
- with_firing_vectors (default: False) boolean

OUTPUT:

list (of divisors)

```
sage: s = sandpiles.Complete(4)
sage: D = SandpileDivisor(s,[4,2,0,0])
sage: D.effective_div()
[{0: 0, 1: 6, 2: 0, 3: 0},
{0: 0, 1: 2, 2: 4, 3: 0},
{0: 0, 1: 2, 2: 0, 3: 4},
{0: 1, 1: 3, 2: 1, 3: 1},
{0: 2, 1: 0, 2: 2, 3: 2},
{0: 4, 1: 2, 2: 0, 3: 0}]
sage: D.effective_div(False)
[[0, 6, 0, 0],
```

```
[0, 2, 4, 0],
 [0, 2, 0, 4],
 [1, 3, 1, 1],
 [2, 0, 2, 2],
 [4, 2, 0, 0]]
sage: D.effective_div(with_firing_vectors=True)
[({0: 0, 1: 6, 2: 0, 3: 0}, (0, -2, -1, -1)),
 (\{0: 0, 1: 2, 2: 4, 3: 0\}, (0, -1, -2, -1)),
 (\{0: 0, 1: 2, 2: 0, 3: 4\}, (0, -1, -1, -2)),
 ({0: 1, 1: 3, 2: 1, 3: 1}, (0, -1, -1, -1)),
 (\{0: 2, 1: 0, 2: 2, 3: 2\}, (0, 0, -1, -1)),
 ({0: 4, 1: 2, 2: 0, 3: 0}, (0, 0, 0, 0))]
sage: a = _[0]
sage: a[0].values()
[0, 6, 0, 0]
sage: vector(D.values()) - s.laplacian()*a[1]
(0, 6, 0, 0)
sage: D.effective_div(False, True)
[([0, 6, 0, 0], (0, -2, -1, -1)),
 ([0, 2, 4, 0], (0, -1, -2, -1)),
 ([0, 2, 0, 4], (0, -1, -1, -2)),
 ([1, 3, 1, 1], (0, -1, -1, -1)),
 ([2, 0, 2, 2], (0, 0, -1, -1)),
 ([4, 2, 0, 0], (0, 0, 0, 0))]
sage: D = SandpileDivisor(s, [-1, 0, 0, 0])
sage: D.effective_div(False,True)
[]
```

— fire_script(sigma)

Fire the given script. In other words, fire each vertex the number of times indicated by sigma.

INPUT:

sigma – SandpileDivisor or (list or dict representing a SandpileDivisor)

OUTPUT:

SandpileDivisor

EXAMPLES:

```
sage: S = sandpiles.Cycle(3)
sage: D = SandpileDivisor(S, [1,2,3])
sage: D.unstable()
[1, 2]
sage: D.fire_script([0,1,1])
{0: 3, 1: 1, 2: 2}
sage: D.fire_script(SandpileDivisor(S,[2,0,0])) == D.fire_vertex(0).fire_vertex(0)
True
```

— fire unstable()

Fire all unstable vertices.

OUTPUT:

SandpileDivisor

```
sage: S = sandpiles.Cycle(3)
sage: D = SandpileDivisor(S, [1,2,3])
sage: D.fire_unstable()
\{0: 3, 1: 1, 2: 2\}
— fire_vertex(v)
Fire the given vertex.
INPUT:
∨ - vertex
OUTPUT:
SandpileDivisor
EXAMPLES:
sage: S = sandpiles.Cycle(3)
sage: D = SandpileDivisor(S, [1,2,3])
sage: D.fire_vertex(1)
{0: 2, 1: 0, 2: 4}
— help(verbose=True)
List of SandpileDivisor methods. If verbose, include short descriptions.
INPUT:
verbose - (default: True) boolean
OUTPUT:
printed string
EXAMPLES:
sage: SandpileDivisor.help()
For detailed help with any method FOO listed below,
enter "SandpileDivisor.FOO?" or enter "D.FOO?" for any SandpileDivisor D.
Dcomplex
                      -- The support-complex.
add_random
                      -- Add one grain of sand to a random vertex.
                      -- The Betti numbers for the support-complex.
betti
                      -- The degree of the divisor.
deg
                     -- The difference with the maximal stable divisor.
dualize
effective_div
                     -- All linearly equivalent effective divisors.
fire_script
                     -- Fire the given script.
fire_unstable
                     -- Fire all unstable vertices.
fire_vertex
                     -- Fire the given vertex.
help
                      -- List of SandpileDivisor methods.
              -- Is the divisor stabilizable?
is alive
is_linearly_equivalent -- Is the given divisor linearly equivalent?
is_q_reduced -- Is the divisor q-reduced?
                      -- Is the divisor symmetric?
is_symmetric
is_weierstrass_pt -- Is the given vertex a Weierstrass point?
linear_system
                      -- The complete linear system (deprecated: use "polytope_integer_pts").
                      -- The polytope determining the complete linear system.
polytope
polytope_integer_pts -- The integer points inside divisor's polytope.
                     -- The linearly equivalent q-reduced divisor.
q_reduced
r_of_D
                      -- The rank of the divisor (deprecated: use "rank", instead).
```

```
-- The rank of the divisor.
rank
                     -- The divisor's underlying sandpile.
sandpile
                     -- Show the divisor.
show
simulate_threshold
                     -- The first unstabilizable divisor in the closed Markov chain.
stabilize
                     -- The stabilization of the divisor.
                     -- List of vertices at which the divisor is nonzero.
support
unstable
                     -- The unstable vertices.
                     -- The values of the divisor as a list.
values
weierstrass_div -- The Weierstrass divisor.
weierstrass_gap_seq -- The Weierstrass gap sequence at the given vertex.
weierstrass_pts -- The Weierstrass points (vertices).
weierstrass_rank_seq -- The Weierstrass rank sequence at the given vertex.
```

— is alive(cycle=False)

Is the divisor stabilizable? In other words, will the divisor stabilize under repeated firings of all unstable vertices? Optionally returns the resulting cycle.

INPUT:

```
cycle - (default: False) boolean
```

OUTPUT:

boolean or optionally, a list of SandpileDivisors

EXAMPLES:

```
sage: S = sandpiles.Complete(4)
sage: D = SandpileDivisor(S, {0: 4, 1: 3, 2: 3, 3: 2})
sage: D.is_alive()
True
sage: D.is_alive(True)
[{0: 4, 1: 3, 2: 3, 3: 2}, {0: 3, 1: 2, 2: 2, 3: 5}, {0: 1, 1: 4, 2: 4, 3: 3}]
```

— is_linearly_equivalent(D, with_firing_vector=False)

Is the given divisor linearly equivalent? Optionally, returns the firing vector. (See NOTE.)

INPUT:

- D SandpileDivisor or list, tuple, etc. representing a divisor
- with_firing_vector (default: False) boolean

OUTPUT:

boolean or integer vector

```
sage: s = sandpiles.Complete(3)
sage: D = SandpileDivisor(s,[2,0,0])
sage: D.is_linearly_equivalent([0,1,1])
True
sage: D.is_linearly_equivalent([0,1,1],True)
(1, 0, 0)
sage: v = vector(D.is_linearly_equivalent([0,1,1],True))
sage: vector(D.values()) - s.laplacian()*v
(0, 1, 1)
sage: D.is_linearly_equivalent([0,0,0])
False
```

```
sage: D.is_linearly_equivalent([0,0,0],True)
()
```

Note:

- If with_firing_vector is False, returns either True or False.
- If with_firing_vector is True then: (i) if self is linearly equivalent to D, returns a vector v such that self v*self.laplacian().transpose() = D. Otherwise, (ii) if self is not linearly equivalent to D, the output is the empty vector, ().

— is_q_reduced()

Is the divisor q-reduced? This would mean that self = c + kq where c is superstable, k is an integer, and q is the sink vertex.

OUTPUT:

boolean

EXAMPLES:

```
sage: s = sandpiles.Complete(4)
sage: D = SandpileDivisor(s,[2,-3,2,0])
sage: D.is_q_reduced()
False
sage: SandpileDivisor(s,[10,0,1,2]).is_q_reduced()
True
```

For undirected or, more generally, Eulerian graphs, q-reduced divisors are linearly equivalent if and only if they are equal. The same does not hold for general directed graphs:

```
sage: s = Sandpile({0:[1],1:[1,1]})
sage: D = SandpileDivisor(s,[-1,1])
sage: Z = s.zero_div()
sage: D.is_q_reduced()
True
sage: Z.is_q_reduced()
True
sage: D == Z
False
sage: D.is_linearly_equivalent(Z)
True
```

— is symmetric(orbits)

Is the divisor symmetric? Return True if the values of the configuration are constant over the vertices in each sublist of orbits.

INPUT:

orbits - list of lists of vertices

OUTPUT:

boolean

```
sage: S = sandpiles.House()
sage: S.dict()
```

```
{0: {1: 1, 2: 1},
1: {0: 1, 3: 1},
 2: {0: 1, 3: 1, 4: 1},
 3: {1: 1, 2: 1, 4: 1},
 4: {2: 1, 3: 1}}
sage: D = SandpileDivisor(S, [0,0,1,1,3])
sage: D.is_symmetric([[2,3], [4]])
True
— is_weierstrass_pt(v='sink')
Is the given vertex a Weierstrass point?
INPUT:
v – (default: sink) vertex
OUTPUT:
boolean
EXAMPLES:
sage: s = sandpiles.House()
sage: K = s.canonical_divisor()
sage: K.weierstrass_rank_seq() # sequence at the sink vertex, 0
(1, 0, -1)
sage: K.is_weierstrass_pt()
False
sage: K.weierstrass_rank_seq(4)
(1, 0, 0, -1)
sage: K.is_weierstrass_pt(4)
```

Note: The vertex v is a (generalized) Weierstrass point for divisor D if the sequence of ranks r(D-nv) for $n=0,1,2,\ldots$ is not $r(D),r(D)-1,\ldots,0,-1,-1,\ldots$

— polytope()

The polytope determining the complete linear system.

OUTPUT:

polytope

EXAMPLES:

```
sage: s = sandpiles.Complete(4)
sage: D = SandpileDivisor(s,[4,2,0,0])
sage: p = D.polytope()
sage: p.inequalities()
(An inequality (-3, 1, 1) x + 2 >= 0,
   An inequality (1, 1, 1) x + 4 >= 0,
   An inequality (1, -3, 1) x + 0 >= 0,
   An inequality (1, 1, -3) x + 0 >= 0)
sage: D = SandpileDivisor(s,[-1,0,0,0])
sage: D.polytope()
The empty polyhedron in QQ^3
```

Note: For a divisor D, this is the intersection of (i) the polyhedron determined by the system of inequalities $L^t x \leq D$

where L^t is the transpose of the Laplacian with (ii) the hyperplane $x_{\text{sink_vertex}} = 0$. The polytope is thought of as sitting in (n-1)-dimensional Euclidean space where n is the number of vertices.

— polytope_integer_pts()

The integer points inside divisor's polytope. The polytope referred to here is the one determining the divisor's complete linear system (see the documentation for polytope).

OUTPUT:

tuple of integer vectors

EXAMPLES:

— q_reduced(verbose=True)

The linearly equivalent q-reduced divisor.

INPUT:

verbose - (default: True) boolean

OUTPUT

SandpileDivisor or list representing SandpileDivisor

EXAMPLES:

```
sage: s = sandpiles.Complete(4)
sage: D = SandpileDivisor(s,[2,-3,2,0])
sage: D.q_reduced()
{0: -2, 1: 1, 2: 2, 3: 0}
sage: D.q_reduced(False)
[-2, 1, 2, 0]
```

Note: The divisor D is qreducedif'D = c + kq where c is superstable, k is an integer, and q is the sink.

— rank(with witness=False)

The rank of the divisor. Optionally returns an effective divisor E such that D-E is not winnable (has an empty complete linear system).

INPUT:

```
with_witness - (default: False) boolean
```

OUTPUT:

integer or (integer, SandpileDivisor)

EXAMPLES:

```
sage: S = sandpiles.Complete(4)
   sage: D = SandpileDivisor(S, [4, 2, 0, 0])
   sage: D.rank()
   sage: D.rank(True)
   (3, \{0: 3, 1: 0, 2: 1, 3: 0\})
   sage: E = [1]
   sage: (D - E).rank()
   -1
Riemann-Roch theorem::
   sage: D.rank() - (S.canonical_divisor()-D).rank() == D.deg() + 1 - S.genus()
Riemann-Roch theorem::
   sage: D.rank() - (S.canonical_divisor()-D).rank() == D.deg() + 1 - S.genus()
   sage: S = Sandpile(\{0:[1,1,1,2],1:[0,0,0,1,1,1,2,2],2:[2,2,1,1,0]\},0) \# multigraph with loops
   sage: D = SandpileDivisor(S, [4, 2, 0])
   sage: D.rank(True)
   (2, \{0: 1, 1: 1, 2: 1\})
   sage: S = Sandpile({0:[1,2], 1:[0,2,2], 2: [0,1]},0) # directed graph
   sage: S.is_undirected()
   False
   sage: D = SandpileDivisor(S, [0, 2, 0])
   sage: D.effective_div()
   [\{0: 0, 1: 2, 2: 0\}, \{0: 2, 1: 0, 2: 0\}]
   sage: D.rank(True)
   (0, \{0: 0, 1: 0, 2: 1\})
   sage: E = D.rank(True)[1]
   sage: (D - E).effective_div()
   []
```

Note: The rank of a divisor D is -1 if D is not linearly equivalent to an effective divisor (i.e., the dollar game represented by D is unwinnable). Otherwise, the rank of D is the largest integer r such that D - E is linearly equivalent to an effective divisor for all effective divisors E with $\deg(E) = r$.

— sandpile()

The divisor's underlying sandpile.

OUTPUT:

Sandpile

EXAMPLES:

```
sage: S = sandpiles.Diamond()
sage: D = SandpileDivisor(S,[1,-2,0,3])
sage: D.sandpile()
Diamond sandpile graph: 4 vertices, sink = 0
sage: D.sandpile() == S
True
```

— show(heights=True, directed=None, **kwds)

Show the divisor.

INPUT:

- heights (default: True) whether to label each vertex with the amount of sand
- directed (optional) whether to draw directed edges
- kwds (optional) arguments passed to the show method for Graph

EXAMPLES:

```
sage: S = sandpiles.Diamond()
sage: D = SandpileDivisor(S,[1,-2,0,2])
sage: D.show(graph_border=True,vertex_size=700,directed=False)
```

— simulate threshold(distrib=None)

The first unstabilizable divisor in the closed Markov chain. (See NOTE.)

INPUT:

distrib – (optional) list of nonnegative numbers representing a probability distribution on the vertices

OUTPUT:

SandpileDivisor

EXAMPLES:

```
sage: s = sandpiles.Complete(4)
sage: D = s.zero_div()
sage: D.simulate_threshold() # random
{0: 2, 1: 3, 2: 1, 3: 2}
sage: n(mean([D.simulate_threshold().deg() for _ in range(10)])) # random
7.10000000000000
sage: n(s.stationary_density()*s.num_verts())
6.93750000000000
```

Note: Starting at self, repeatedly choose a vertex and add a grain of sand to it. Return the first unstabilizable divisor that is reached. Also see the markov_chain method for the underlying sandpile.

— stabilize(with_firing_vector=False)

The stabilization of the divisor. If not stabilizable, return an error.

INPUT:

```
with_firing_vector - (default: False) boolean
```

EXAMPLES:

```
sage: s = sandpiles.Complete(4)
sage: D = SandpileDivisor(s,[0,3,0,0])
sage: D.stabilize()
{0: 1, 1: 0, 2: 1, 3: 1}
sage: D.stabilize(with_firing_vector=True)
[{0: 1, 1: 0, 2: 1, 3: 1}, {0: 0, 1: 1, 2: 0, 3: 0}]
```

— support()

List of vertices at which the divisor is nonzero.

OUTPUT:

list representing the support of the divisor

EXAMPLES:

```
sage: S = sandpiles.Cycle(4)
sage: D = SandpileDivisor(S, [0,0,1,1])
sage: D.support()
[2, 3]
sage: S.vertices()
[0, 1, 2, 3]
```

— unstable()

The unstable vertices.

OUTPUT:

list of vertices

EXAMPLES:

```
sage: S = sandpiles.Cycle(3)
sage: D = SandpileDivisor(S, [1,2,3])
sage: D.unstable()
[1, 2]
```

— values()

The values of the divisor as a list. The list is sorted in the order of the vertices.

OUTPUT:

list of integers

boolean

EXAMPLES:

```
sage: S = Sandpile({'a':[1,'b'], 'b':[1,'a'], 1:['a']},'a')
sage: D = SandpileDivisor(S, {'a':0, 'b':1, 1:2})
sage: D
{'a': 0, 1: 2, 'b': 1}
sage: D.values()
[2, 0, 1]
sage: S.vertices()
[1, 'a', 'b']
```

— weierstrass_div(verbose=True)

The Weierstrass divisor. Its value at a vertex is the weight of that vertex as a Weierstrass point. (See SandpileDivisor.weierstrass_gap_seq.)

INPUT:

verbose - (default: True) boolean

OUTPUT:

SandpileDivisor

```
sage: s = sandpiles.Diamond()
sage: D = SandpileDivisor(s,[4,2,1,0])
sage: [D.weierstrass_rank_seq(v) for v in s]
[(5, 4, 3, 2, 1, 0, 0, -1),
  (5, 4, 3, 2, 1, 0, 0, 0, -1),
  (5, 4, 3, 2, 1, 0, 0, 0, -1),
  (5, 4, 3, 2, 1, 0, 0, -1)]
sage: D.weierstrass_div()
{0: 1, 1: 0, 2: 2, 3: 1}
sage: k5 = sandpiles.Complete(5)
sage: K = k5.canonical_divisor()
sage: K.weierstrass_div()
{0: 9, 1: 9, 2: 9, 3: 9, 4: 9}
```

— weierstrass_gap_seq(v='sink', weight=True)

The Weierstrass gap sequence at the given vertex. If weight is True, then also compute the weight of each gap value.

INPUT:

- v (default: sink) vertex
- weight (default: True) boolean

OUTPUT:

list or (list of list) of integers

EXAMPLES:

```
sage: s = sandpiles.Cycle(4)
sage: D = SandpileDivisor(s,[2,0,0,0])
sage: [D.weierstrass_gap_seq(v,False) for v in s.vertices()]
[(1, 3), (1, 2), (1, 3), (1, 2)]
sage: [D.weierstrass_gap_seq(v) for v in s.vertices()]
[((1, 3), 1), ((1, 2), 0), ((1, 3), 1), ((1, 2), 0)]
sage: D.weierstrass_gap_seq() # gap sequence at sink vertex, 0
((1, 3), 1)
sage: D.weierstrass_rank_seq() # rank sequence at the sink vertex
(1, 0, 0, -1)
```

Note: The integer k is a Weierstrass gap for the divisor D at vertex v if the rank of D-(k-1)v does not equal the rank of D-kv. Let r be the rank of D and let k_i be the i-th gap at v. The Weierstrass weight of v for D is the sum of (k_i-i) as i ranges from 1 to r+1. It measure the difference between the sequence r, r-1, ..., 0, -1, -1, ... and the rank sequence $\operatorname{rank}(D), \operatorname{rank}(D-v), \operatorname{rank}(D-2v), ...$

— weierstrass_pts(with_rank_seq=False)

The Weierstrass points (vertices). Optionally, return the corresponding rank sequences.

INPUT:

```
with_rank_seq - (default: False) boolean
```

OUTPUT:

tuple of vertices or list of (vertex, rank sequence)

EXAMPLES:

```
sage: s = sandpiles.House()
sage: K = s.canonical_divisor()
sage: K.weierstrass_pts()
(4,)
sage: K.weierstrass_pts(True)
[(4, (1, 0, 0, -1))]
```

Note: The vertex v is a (generalized) Weierstrass point for divisor D if the sequence of ranks r(D-nv) for $n=0,1,2,\ldots$ 'is not $r(D),r(D)-1,\ldots,0,-1,-1,\ldots$

— weierstrass_rank_seq(v='sink')

The Weierstrass rank sequence at the given vertex. Computes the rank of the divisor D - nv starting with n = 0 and ending when the rank is -1.

INPUT:

```
v – (default: sink) vertex
```

OUTPUT:

tuple of int

EXAMPLES:

```
sage: s = sandpiles.House()
sage: K = s.canonical_divisor()
sage: [K.weierstrass_rank_seq(v) for v in s.vertices()]
[(1, 0, -1), (1, 0, -1), (1, 0, -1), (1, 0, -1), (1, 0, 0, -1)]
```

Other

- firing_graph The firing graph.
- *parallel_firing_graph* The parallel-firing graph.
- random_DAG A random directed acyclic graph.
- sandpiles Some examples of sandpiles.
- wilmes_algorithm Find matrix with the same integer row span as M that is the reduced Laplacian of a digraph.

Complete descriptions of methods. firing_graph(S, eff)

Creates a digraph with divisors as vertices and edges between two divisors D and E if firing a single vertex in D gives E.

INPUT:

S - Sandpile eff - list of divisors

OUTPUT:

DiGraph

EXAMPLES:

```
sage: S = sandpiles.Cycle(6)
sage: D = SandpileDivisor(S, [1,1,1,1,2,0])
sage: eff = D.effective_div()
sage: firing_graph(S,eff).show3d(edge_size=.005,vertex_size=0.01)
```

— parallel_firing_graph(S,eff)

Creates a digraph with divisors as vertices and edges between two divisors D and E if firing all unstable vertices in D gives E.

INPUT:

S - Sandpile eff - list of divisors

OUTPUT:

DiGraph

EXAMPLES:

```
sage: S = Sandpile(graphs.CycleGraph(6),0)
sage: D = SandpileDivisor(S, [1,1,1,1,2,0])
sage: eff = D.effective_div()
sage: parallel_firing_graph(S,eff).show3d(edge_size=.005,vertex_size=0.01)
```

— random_DAG(num_verts,p=1/2,weight_max=1)

Returns a random directed acyclic graph with num_verts vertices. The method starts with the sink vertex and adds vertices one at a time. Each vertex is connected only to only previously defined vertices, and the probability of each possible connection is given by the argument p. The weight of an edge is a random integer between 1 and weight_max.

INPUT:

- num_verts positive integer
- p number between 0 and 1
- weight_max integer greater than 0

OUTPUT:

directed acyclic graph with sink 0

EXAMPLES:

```
sage: S = random_DAG(5, 0.3)
```

- sandpiles

Some examples of sandpiles.

Here are the available examples; you can also type "sandpiles." and hit tab to get a list:

- "Complete()"
- "Cycle()"
- "Diamond()"
- "Grid()"
- "House()"

EXAMPLES:

```
sage: s = sandpiles.Complete(4)
sage: s.invariant_factors()
[1, 4, 4]
sage: s.laplacian()
[ 3 -1 -1 -1]
[-1  3 -1 -1]
[-1 -1  3 -1]
[-1 -1  3 -1]
```

— wilmes_algorithm(M)

Computes an integer matrix L with the same integer row span as M and such that L is the reduced laplacian of a directed multigraph.

INPUT:

M - square integer matrix of full rank

OUTPUT:

L - integer matrix

EXAMPLES:

NOTES:

The algorithm is due to John Wilmes.

Help

Documentation for each method is available through the Sage online help system:

```
sage: SandpileConfig.fire_vertex?
Base Class: <type 'instancemethod'>
String Form:
               <unbound method SandpileConfig.fire_vertex>
Namespace: Interactive
               /usr/local/sage-4.7/local/lib/python2.6/site-packages/sage/sandpiles/sandpile.py
File:
Definition:
              SandpileConfig.fire_vertex(self, v)
Docstring:
      Fire the vertex ''v''.
       INPUT:
       ''v'' - vertex
       OUTPUT:
       SandpileConfig
       EXAMPLES:
          sage: S = Sandpile(graphs.CycleGraph(3), 0)
          sage: c = SandpileConfig(S, [1,2])
```

```
sage: c.fire_vertex(2)
{1: 2, 2: 0}
```

Note: An alternative to SandpileConfig.fire_vertex? in the preceding code example would be c.fire_vertex?, if c is any SandpileConfig.

Enter Sandpile.help(), SandpileConfig.help(), and SandpileDivisor.help() for lists of available Sandpile-specific methods.

General Sage documentation can be found at http://sagemath.org/doc/.

Contact

Please contact davidp@reed.edu with questions, bug reports, and suggestions for additional features and other improvements.

12.1.4 Group Theory and Sage

Author: Robert A. Beezer, University of Puget Sound

This compilation collects Sage commands that are useful for a student in an introductory course on group theory. It is not intended to teach Sage or to teach group theory. (There are many introductory texts on group theory and more information on Sage can be found via www.sagemath.org) Rather, by presenting commands roughly in the order a student would learn the corresponding mathematics they might be encouraged to experiment and learn more about mathematics and learn more about Sage. Not coincidentally, when Sage was the acronym SAGE, the "E" in Sage stood for "Experimentation."

This guide is also distributed in PDF format and as a Sage worksheet. The worksheet version can be imported into the Sage notebook environment running in a web browser, and then the displayed chunks of code may be executed by Sage if one clicks on the small "evaluate" link below each cell, for a fully interactive experience. A PDF and Sage worksheet versions of this tutorial are available at http://abstract.ups.edu/sage-aata.html.

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- · Group Theory and Sage
 - Basic properties of the integers
 - Permutation groups
 - Group functions
 - Subgroups
 - Symmetry groups
 - Normal subgroups
 - Conjugacy
 - Sylow subgroups
 - Groups of small order as permutation groups
 - Acknowledgements

Changelog:

- 2009/01/30 Version 1.0, first complete release
- 2009/03/03 Version 1.1, added cyclic group size interact
- 2010/03/10 Version 1.3, dropped US on license, some edits.

Basic properties of the integers

Integer division

The command a % b will return the remainder upon division of a by b. In other words, the value is the unique integer r such that:

- 1. $0 \le r \le b$; and
- 2. a = bq + r for some integer q (the quotient).

Then (a - r)/b will equal q. For example:

```
sage: r = 14 % 3
sage: q = (14 - r) / 3
sage: r, q
(2, 4)
```

will return 2 for the value of r and 4 for the value of q. Note that the "/" is *integer* division, where any remainder is cast away and the result is always an integer. So, for example, 14 / 3 will again equal 4, not 4.66666.

Greatest common divisor

The greatest common divisor of a and b is obtained with the command gcd(a,b), where in our first uses, a and b are integers. Later, a and b can be other objects with a notion of divisibility and "greatness," such as polynomials. For example:

```
sage: gcd(2776, 2452)
4
```

Extended greatest common divisor

The command xgcd(a, b) ("eXtended GCD") returns a triple where the first element is the greatest common divisor of a and b (as with the gcd(a, b) command above), but the next two elements are the values of r and s such that ra + sb = gcd(a, b). For example, xgcd(633, 331) returns (1, 194, -371). Portions of the triple can be extracted using [] to access the entries of the triple, starting with the first as number 0. For example, the following should return the result True (even if you change the values of a and b). Studying this block of code will go a long way towards helping you get the most out of Sage's output. (Note that "=" is how a value is assigned to a variable, while as in the last line, "==" is how we determine equality of two items.)

```
sage: a = 633
sage: b = 331
sage: extended = xgcd(a, b)
sage: g = extended[0]
sage: r = extended[1]
sage: s = extended[2]
sage: g == r*a + s*b
True
```

Divisibility

A remainder of zero indicates divisibility. So (a % b) == 0 will return True if b divides a, and will otherwise return False. For example, (9 % 3) == 0 is True, but (9 % 4) == 0 is False. Try predicting the output of the following before executing it in Sage.

```
sage: answer1 = ((20 % 5) == 0)
sage: answer2 = ((17 % 4) == 0)
sage: answer1, answer2
(True, False)
```

Factoring

As promised by the Fundamental Theorem of Arithmetic, factor (a) will return a unique expression for a as a product of powers of primes. It will print in a nicely-readable form, but can also be manipulated with Python as a list of pairs (p_i, e_i) containing primes as bases, and their associated exponents. For example:

```
sage: factor(2600)
2^3 * 5^2 * 13
```

If you just want the prime divisors of an integer, then use the $prime_divisors$ (a) command, which will return a list of all the prime divisors of a. For example:

```
sage: prime_divisors(2600)
[2, 5, 13]
```

We can strip off other pieces of the prime decomposition using two levels of []. This is another good example to study in order to learn about how to drill down into Python lists.

```
sage: n = 2600
sage: decomposition = factor(n)
sage: print n, "decomposes as", decomposition
2600 decomposes as 2^3 * 5^2 * 13
sage: secondterm = decomposition[1]
sage: print "Base and exponent (pair) for second prime:", secondterm
Base and exponent (pair) for second prime: (5, 2)
sage: base = secondterm[0]
sage: exponent = secondterm[1]
sage: print "Base is", base
Base is 5
sage: print "Exponent is", exponent
Exponent is 2
sage: thirdbase = decomposition[2][0]
sage: thirdexponent = decomposition[2][1]
sage: print "Base of third term is", thirdbase, "with exponent", thirdexponent
Base of third term is 13 with exponent 1
```

With a bit more work, the factor () command can be used to factor more complicated items, such as polynomials.

Multiplicative inverse, modular arithmetic

The command inverse_mod(a, n) yields the multiplicative inverse of $a \mod n$ (or an error if it doesn't exist). For example:

```
sage: inverse_mod(352, 917)
508
```

(As a check, find the integer m such that 352*508 = m*917+1.) Then try

```
sage: inverse_mod(4, 24)
Traceback (most recent call last):
...
ZeroDivisionError: Inverse does not exist.
and explain the result.
```

Powers with modular arithmetic

The command power_mod (a, m, n) yields $a^m \mod n$. For example:

```
sage: power_mod(15, 831, 23)
10
```

If m = -1, then this command will duplicate the function of inverse_mod().

Euler ϕ -function

The command $euler_phi(n)$ will return the number of positive integers less than n and relatively prime to n (i.e. having greatest common divisor with n equal to 1). For example:

```
sage: euler_phi(345)
176
```

sage: is_prime(117371)

Experiment by running the following code several times:

```
sage: m = random_prime(10000)
sage: n = random_prime(10000)
sage: euler_phi(m*n) == euler_phi(m) * euler_phi(n)
True
```

Feel a conjecture coming on? Can you generalize this result?

Primes

The command is_prime (a) returns True or False depending on if a is prime or not. For example,

The command random_prime (a, True) will return a random prime between 2 and a. Experiment with:

```
sage: p = random_prime(10^21, True)
sage: is_prime(p)
True
```

(Replacing True by False will speed up the search, but there will be a very small probability the result will not be prime.)

The command prime_range(a, b) returns an ordered list of all the primes from a to b-1, inclusive. For example,

```
sage: prime_range(500, 550)
[503, 509, 521, 523, 541, 547]
```

The commands next_prime (a) and previous_prime (a) are other ways to get a single prime number of a desired size. Give them a try.

Permutation groups

A good portion of Sage's support for group theory is based on routines from GAP (Groups, Algorithms, and Programming at http://www.gap-system.org. Groups can be described in many different ways, such as sets of matrices or sets of symbols subject to a few defining relations. A very concrete way to represent groups is via permutations (one-to-one and onto functions of the integers 1 through n), using function composition as the operation in the group. Sage has many routines designed to work with groups of this type and they are also a good way for those learning group theory to gain experience with the basic ideas of group theory. For both these reasons, we will concentrate on these types of groups.

Writing permutations

Sage uses "disjoint cycle notation" for permutations, see any introductory text on group theory (such as Judson, Section 4.1) for more on this. Composition occurs *left to right*, which is not what you might expect and is exactly the reverse of what Judson and many others use. (There are good reasons to support either direction, you just need to be certain you know which one is in play.) There are two ways to write the permutation $\sigma = (13)(254)$:

- 1. As a text string (include quotes): " (1, 3) (2, 5, 4) "
- 2. As a Python list of "tuples": [(1,3), (2,5,4)]

Groups

Sage knows many popular groups as sets of permutations. More are listed below, but for starters, the full "symmetric group" of all possible permutations of 1 through n can be built with the command SymmetricGroup (n).

Permutation elements Elements of a group can be created, and composed, as follows

```
sage: G = SymmetricGroup(5)
sage: sigma = G("(1,3) (2,5,4)")
sage: rho = G([(2,4), (1,5)])
sage: rho^(-1) * sigma * rho
(1,2,4)(3,5)
```

Available functions for elements of a permutation group include finding the order of an element, i.e. for a permutation σ the order is the smallest power of k such that σ^k equals the identity element (). For example:

```
sage: G = SymmetricGroup(5)
sage: sigma = G("(1,3) (2,5,4)")
sage: sigma.order()
```

The sign of the permutation σ is defined to be 1 for an even permutation and -1 for an odd permutation. For example:

```
sage: G = SymmetricGroup(5)
sage: sigma = G("(1,3) (2,5,4)")
sage: sigma.sign()
-1
```

since σ is an odd permutation.

Many more available functions that can be applied to a permutation can be found via "tab-completion." With sigma defined as an element of a permutation group, in a Sage cell, type sigma. (Note the ".") and then press the tab key. You will get a list of available functions (you may need to scroll down to see the whole list). Experiment and explore! It is what Sage is all about. You really cannot break anything.

Creating groups

This is an annotated list of some small well-known permutation groups that can be created simply in Sage. You can find more in the source code file

SAGE_ROOT/src/sage/groups/perm_gps/permgroup_named.py

- SymmetricGroup (n): All n! permutations on n symbols.
- DihedralGroup (n): Symmetries of an n-gon. Rotations and flips, 2n in total.
- CyclicPermutationGroup (n): Rotations of an n-gon (no flips), n in total.
- Alternating Group (n): Alternating group on n symbols having n!/2 elements.
- KleinFourGroup (): The non-cyclic group of order 4.

Group functions

Individual elements of permutation groups are important, but we primarily wish to study groups as objects on their own. So a wide variety of computations are available for groups. Define a group, for example

```
sage: H = DihedralGroup(6)
sage: H
Dihedral group of order 12 as a permutation group
```

and then a variety of functions become available.

After trying the examples below, experiment with tab-completion. Having defined H, type H. (note the ".") and then press the tab key. You will get a list of available functions (you may need to scroll down to see the whole list). As before, *experiment and explore*—it is really hard to break anything.

Here is another couple of ways to experiment and explore. Find a function that looks interesting, say is_abelian(). Type H.is_abelian? (note the question mark) followed by the enter key. This will display a portion of the source code for the is_abelian() function, describing the inputs and output, possibly illustrated with example uses.

If you want to learn more about how Sage works, or possibly extend its functionality, then you can start by examining the complete Python source code. For example, try <code>H.is_abelian??</code>, which will allow you to determine that the <code>is_abelian()</code> function is basically riding on GAP's <code>IsAbelian()</code> command and asking GAP do the heavy-lifting for us. (To get the maximum advantage of using Sage it helps to know some basic Python programming, but it is not required.)

OK, on to some popular command for groups. If you are using the worksheet, be sure you have defined the group H as the dihedral group D_6 , since we will not keep repeating its definition below.

Abelian?

The command

```
sage: H = DihedralGroup(6)
sage: H.is_abelian()
False
```

will return False since D_6 is a non-abelian group.

Order

The command

```
sage: H = DihedralGroup(6)
sage: H.order()
12
```

will return 12 since D_6 is a group of with 12 elements.

All elements

The command

```
sage: H = DihedralGroup(6)
sage: H.list()
[(),
    (1,6)(2,5)(3,4),
    (1,2,3,4,5,6),
    (1,5)(2,4),
    (2,6)(3,5),
    (1,3,5)(2,4,6),
    (1,4)(2,3)(5,6),
    (1,6,5,4,3,2),
    (1,4)(2,5)(3,6),
    (1,2)(3,6)(4,5),
    (1,5,3)(2,6,4),
    (1,3)(4,6)]
```

will return all of the elements of H in a fixed order as a Python list. Indexing ($[\]$) can be used to extract the individual elements of the list, remembering that counting the elements of the list begins at zero.

```
sage: H = DihedralGroup(6)
sage: elements = H.list()
sage: elements[2]
(1,2,3,4,5,6)
```

Cayley table

The command

```
sage: H = DihedralGroup(6)
sage: H.cayley_table()
* a b c d e f g h i j k l
```

```
a| a b c d e f g h i j k l
b| b a e h c j k d l f g i
c| c d f g b i l a k e h j
d| d c b a f e h g j i l k
e| e h j k a l i b g c d f
f| f g i l d k j c h b a e
g| g f d c i b a l e k j h
h| h e a b j c d k f l i g
i| i l k j g h e f a d c b
j| j k l i h g f e d a b c
k| k j h e l a b i c g f d
l l i g f k d c j b h e a
```

will construct the Cayley table (or "multiplication table") of H. By default the table uses lowercase Latin letters to name the elements of the group. The actual elements used can be found using the row_keys() or column_keys() commands for the table. For example to determine the fifth element in the table, the element named e:

```
sage: H = DihedralGroup(6)
sage: T = H.cayley_table()
sage: headings = T.row_keys()
sage: headings[4]
(2,6)(3,5)
```

Center

The command ${\tt H.center}$ () will return a subgroup that is the center of the group H (see Exercise 2.46 in Judson). Try

```
sage: H = DihedralGroup(6)
sage: H.center().list()
[(), (1,4)(2,5)(3,6)]
```

to see which elements of H commute with *every* element of H.

Cayley graph

For fun, try show (H.cayley_graph()).

Subgroups

Cyclic subgroups

If G is a group and a is an element of the group (try $a = G.random_element()$), then

```
a = G.random_element()
H = G.subgroup([a])
```

will create H as the cyclic subgroup of G with generator a.

For example the code below will:

1. create G as the symmetric group on five symbols;

- 2. specify sigma as an element of G;
- 3. use sigma as the generator of a cyclic subgroup H;
- 4. list all the elements of H.

In more mathematical notation, we might write $\langle (1\,2\,3)(4\,5) \rangle = H \subseteq G = S_5$.

```
sage: G = SymmetricGroup(5)
sage: sigma = G("(1,2,3) (4,5)")
sage: H = G.subgroup([sigma])
sage: H.list()
[(), (1,2,3)(4,5), (1,3,2), (4,5), (1,2,3), (1,3,2)(4,5)]
```

Experiment by trying different permutations for sigma and observing the effect on H.

Cyclic groups

Groups that are cyclic themselves are both important and rich in structure. The command CyclicPermutationGroup(n) will create a permutation group that is cyclic with n elements. Consider the following example (note that the indentation of the third line is critical) which will list the elements of a cyclic group of order 20, preceded by the order of each element.

```
sage: n = 20
sage: CN = CyclicPermutationGroup(n)
sage: for g in CN:
           print g.order(), " ", g
1
2.0
       (1,2,3,4,5,6,7,8,9,10,11,12,13,14,15,16,17,18,19,20)
10
       (1,3,5,7,9,11,13,15,17,19) (2,4,6,8,10,12,14,16,18,20)
       (1, 4, 7, 10, 13, 16, 19, 2, 5, 8, 11, 14, 17, 20, 3, 6, 9, 12, 15, 18)
2.0
      (1, 5, 9, 13, 17) (2, 6, 10, 14, 18) (3, 7, 11, 15, 19) (4, 8, 12, 16, 20)
4
      (1,6,11,16) (2,7,12,17) (3,8,13,18) (4,9,14,19) (5,10,15,20)
10
      (1,7,13,19,5,11,17,3,9,15) (2,8,14,20,6,12,18,4,10,16)
       (1, 8, 15, 2, 9, 16, 3, 10, 17, 4, 11, 18, 5, 12, 19, 6, 13, 20, 7, 14)
20
5
     (1, 9, 17, 5, 13) (2, 10, 18, 6, 14) (3, 11, 19, 7, 15) (4, 12, 20, 8, 16)
20
      (1,10,19,8,17,6,15,4,13,2,11,20,9,18,7,16,5,14,3,12)
2
     (1,11)(2,12)(3,13)(4,14)(5,15)(6,16)(7,17)(8,18)(9,19)(10,20)
20
      (1, 12, 3, 14, 5, 16, 7, 18, 9, 20, 11, 2, 13, 4, 15, 6, 17, 8, 19, 10)
5
     (1,13,5,17,9) (2,14,6,18,10) (3,15,7,19,11) (4,16,8,20,12)
20
      (1,14,7,20,13,6,19,12,5,18,11,4,17,10,3,16,9,2,15,8)
1.0
      (1, 15, 9, 3, 17, 11, 5, 19, 13, 7) (2, 16, 10, 4, 18, 12, 6, 20, 14, 8)
4
     (1,16,11,6) (2,17,12,7) (3,18,13,8) (4,19,14,9) (5,20,15,10)
     (1,17,13,9,5) (2,18,14,10,6) (3,19,15,11,7) (4,20,16,12,8)
20
      (1,18,15,12,9,6,3,20,17,14,11,8,5,2,19,16,13,10,7,4)
       (1,19,17,15,13,11,9,7,5,3) (2,20,18,16,14,12,10,8,6,4)
10
       (1, 20, 19, 18, 17, 16, 15, 14, 13, 12, 11, 10, 9, 8, 7, 6, 5, 4, 3, 2)
```

By varying the size of the group (change the value of n) you can begin to illustrate some of the structure of a cyclic group (for example, try a prime).

We can cut/paste an element of order 5 from the output above (in the case when the cyclic group has 20 elements) and quickly build a subgroup:

```
sage: C20 = CyclicPermutationGroup(20)
sage: rho = C20("(1,17,13,9,5)(2,18,14,10,6)(3,19,15,11,7)(4,20,16,12,8)")
sage: H = C20.subgroup([rho])
```

```
sage: H.list()
[(),
    (1,17,13,9,5) (2,18,14,10,6) (3,19,15,11,7) (4,20,16,12,8),
    (1,13,5,17,9) (2,14,6,18,10) (3,15,7,19,11) (4,16,8,20,12),
    (1,9,17,5,13) (2,10,18,6,14) (3,11,19,7,15) (4,12,20,8,16),
    (1,5,9,13,17) (2,6,10,14,18) (3,7,11,15,19) (4,8,12,16,20)]
```

For a cyclic group, the following command will list *all* of the subgroups.

```
sage: C20 = CyclicPermutationGroup(20)
sage: C20.conjugacy_classes_subgroups()
[Subgroup of (Cyclic group of order 20 as a permutation group) generated by [()], Subgroup of (Cyclic
```

Be careful, this command uses some more advanced ideas and will not usually list *all* of the subgroups of a group. Here we are relying on special properties of cyclic groups (but see the next section).

If you are viewing this as a PDF, you can safely skip over the next bit of code. However, if you are viewing this as a worksheet in Sage, then this is a place where you can experiment with the structure of the subgroups of a cyclic group. In the input box, enter the order of a cyclic group (numbers between 1 and 40 are good initial choices) and Sage will list each subgroup as a cyclic group with its generator. The factorization at the bottom might help you formulate a conjecture.

All subgroups

If H is a subgroup of G and $g \in G$, then $gHg^{-1} = \{ghg^{-1} \mid h \in G\}$ will also be a subgroup of G. If G is a group, then the command G.conjugacy_classes_subgroups () will return a list of subgroups of G, but not all of the subgroups. However, every subgroup can be constructed from one on the list by the gHg^{-1} construction with a suitable g. As an illustration, the code below:

- 1. creates K as the dihedral group of order 24, D_{12} ;
- 2. stores the list of subgroups output by K. conjugacy_classes_subgroups() in the variable sq;
- 3. prints the elements of the list;
- 4. selects the second subgroup in the list, and lists its elements.

```
sage: K = DihedralGroup(12)
sage: sg = K.conjugacy_classes_subgroups()
sage: print "sg:\n", sg
sg:
[Subgroup of (Dihedral group of order 24 as a permutation group) generated by [()], Subgroup of (Dihedral group of (Dihedral group of (Dihedral group))
```

```
sage: print "\nAn order two subgroup:\n", sg[1].list()
An order two subgroup:
[(), (1,2)(3,12)(4,11)(5,10)(6,9)(7,8)]
```

It is important to note that this is a nice long list of subgroups, but will rarely create *every* such subgroup. For example, the code below:

- 1. creates rho as an element of the group K;
- 2. creates L as a cyclic subgroup of K;
- 3. prints the two elements of L; and finally
- 4. tests to see if this subgroup is part of the output of the list sq created just above (it is not).

```
sage: K = DihedralGroup(12)
sage: sg = K.conjugacy_classes_subgroups()
sage: rho = K("(1,4) (2,3) (5,12) (6,11) (7,10) (8,9)")
sage: L = PermutationGroup([rho])
sage: L.list()
[(), (1,4)(2,3)(5,12)(6,11)(7,10)(8,9)]
sage: L in sg
False
```

Symmetry groups

You can give Sage a short list of elements of a permutation group and Sage will find the smallest subgroup that contains those elements. We say the list "generates" the subgroup. We list a few interesting subgroups you can create this way.

Symmetries of an equilateral triangle

Label the vertices of an equilateral triangle as 1, 2 and 3. Then *any* permutation of the vertices will be a symmetry of the triangle. So either SymmetricGroup (3) or DihedralGroup (3) will create the full symmetry group.

Symmetries of an n-gon

A regular, n-sided figure in the plane (an n-gon) has 2n symmetries, comprised of n rotations (including the trivial one) and n "flips" about various axes. The dihedral group <code>DihedralGroup(n)</code> is frequently defined as exactly the symmetry group of an n-gon.

Symmetries of a tetrahedron

Label the 4 vertices of a regular tetrahedron as 1, 2, 3 and 4. Fix the vertex labeled 4 and rotate the opposite face through 120 degrees. This will create the permutation/symmetry $(1\,2\,3)$. Similarly, fixing vertex 1, and rotating the opposite face will create the permutation $(2\,3\,4)$. These two permutations are enough to generate the full group of the twelve symmetries of the tetrahedron. Another symmetry can be visualized by running an axis through the midpoint of an edge of the tetrahedron through to the midpoint of the opposite edge, and then rotating by 180 degrees about this axis. For example, the 1–2 edge is opposite the 3–4 edge, and the symmetry is described by the permutation $(1\,2)(3\,4)$. This permutation, along with either of the above permutations will also generate the group. So here are two ways to create this group:

```
sage: tetra_one = PermutationGroup(["(1,2,3)", "(2,3,4)"])
sage: tetra_one
Permutation Group with generators [(2,3,4), (1,2,3)]
sage: tetra_two = PermutationGroup(["(1,2,3)", "(1,2)(3,4)"])
sage: tetra_two
Permutation Group with generators [(1,2)(3,4), (1,2,3)]
```

This group has a variety of interesting properties, so it is worth experimenting with. You may also know it as the "alternating group on 4 symbols," which Sage will create with the command AlternatingGroup (4).

Symmetries of a cube

Label vertices of one face of a cube with 1, 2, 3 and 4, and on the opposite face label the vertices 5, 6, 7 and 8 (5 opposite 1, 6 opposite 2, etc.). Consider three axes that run from the center of a face to the center of the opposite face, and consider a quarter-turn rotation about each axis. These three rotations will construct the entire symmetry group. Use

```
sage: cube = PermutationGroup(["(3,2,6,7)(4,1,5,8)",
          "(1,2,6,5)(4,3,7,8)", "(1,2,3,4)(5,6,7,8)"])
sage: cube.list()
[(),
 (1,2,3,4)(5,6,7,8),
 (1,2,6,5)(3,7,8,4),
 (1,5,8,4)(2,6,7,3),
 (1,6,8)(2,7,4),
 (1,3,8)(2,7,5),
 (1,6,3)(4,5,7),
 (1,6)(2,5)(3,8)(4,7),
 (2,5,4)(3,6,8),
 (1,3)(2,4)(5,7)(6,8),
 (1,8)(2,7)(3,6)(4,5),
 (1,7)(2,3)(4,6)(5,8),
 (1,5,6,2)(3,4,8,7),
 (1,7)(2,6)(3,5)(4,8),
 (1,7)(2,8)(3,4)(5,6),
 (1,4,3,2)(5,8,7,6),
 (1,4)(2,8)(3,5)(6,7),
 (1,5)(2,8)(3,7)(4,6),
 (1,4,8,5)(2,3,7,6),
 (1,2)(3,5)(4,6)(7,8),
 (1,8,6)(2,4,7),
 (1,3,6)(4,7,5),
 (2,4,5)(3,8,6),
 (1,8,3)(2,5,7)
```

A cube has four distinct diagonals (joining opposite vertices through the center of the cube). Each symmetry of the cube will cause the diagonals to arrange differently. In this way, we can view an element of the symmetry group as a permutation of four "symbols"—the diagonals. It happens that *each* of the 24 permutations of the diagonals is created by exactly one symmetry of the 8 vertices of the cube. So this subgroup of S_8 is "the same as" S_4 . In Sage:

```
sage: cube = PermutationGroup(["(3,2,6,7)(4,1,5,8)",
...: "(1,2,6,5)(4,3,7,8)", "(1,2,3,4)(5,6,7,8)"])
sage: cube.is_isomorphic(SymmetricGroup(4))
```

will test to see if the group of symmetries of the cube are "the same as" S_4 and so will return True.

Here is an another way to create the symmetries of a cube. Number the six *faces* of the cube as follows: 1 on top, 2 on the bottom, 3 in front, 4 on the right, 5 in back, 6 on the left. Now the same rotations as before (quarter-turns about axes through the centers of two opposite faces) can be used as generators of the symmetry group:

```
sage: cubeface = PermutationGroup(["(1,3,2,5)", "(1,4,2,6)", "(3,4,5,6)"])
sage: cubeface.list()
[(),
 (3,4,5,6),
 (1, 4, 2, 6),
 (1,3,2,5),
 (1,3,4)(2,5,6),
 (1,3,6)(2,5,4),
 (1,2)(3,5),
 (1,4,5)(2,6,3),
 (1,5,6)(2,3,4),
 (3,5)(4,6),
 (1,2)(4,6),
 (1, 5, 2, 3),
 (1,6)(2,4)(3,5),
 (1,4)(2,6)(3,5),
 (1,2)(3,4)(5,6),
 (1,3)(2,5)(4,6),
 (3,6,5,4),
 (1,5)(2,3)(4,6),
 (1, 6, 2, 4),
 (1,2)(3,6)(4,5),
 (1,6,3)(2,4,5),
 (1,6,5)(2,4,3),
 (1,5,4)(2,3,6),
 (1,4,3)(2,6,5)
```

Again, this subgroup of S_6 is "same as" the full symmetric group, S_4 :

```
sage: cubeface = PermutationGroup(["(1,3,2,5)", "(1,4,2,6)", "(3,4,5,6)"])
sage: cubeface.is_isomorphic(SymmetricGroup(4))
True
```

It turns out that in each of the above constructions, it is sufficient to use just two of the three generators (any two). But one generator is not enough. Give it a try and use Sage to convince yourself that a generator can be sacrificed in each case.

Normal subgroups

Checking normality

The code below:

- 1. begins with the alternating group A_4 ;
- 2. specifies three elements of the group (the three symmetries of the tetrahedron that are 180 degree rotations about axes through midpoints of opposite edges);
- 3. uses these three elements to generate a subgroup; and finally
- 4. illustrates the command for testing if the subgroup H is a normal subgroup of the group A4.

```
sage: A4 = AlternatingGroup(4)
sage: r1 = A4("(1,2) (3,4)")
```

```
sage: r2 = A4("(1,3) (2,4)")
sage: r3 = A4("(1,4) (2,3)")
sage: H = A4.subgroup([r1, r2, r3])
sage: H.is_normal(A4)
True
```

Quotient group

Extending the previous example, we can create the quotient (factor) group of A_4 by H. The commands

```
sage: A4 = AlternatingGroup(4)
sage: r1 = A4("(1,2) (3,4)")
sage: r2 = A4("(1,3) (2,4)")
sage: r3 = A4("(1,4) (2,3)")
sage: H = A4.subgroup([r1, r2, r3])
sage: A4.quotient(H)
Permutation Group with generators [(1,2,3)]
```

returns a permutation group generated by (1,2,3). As expected this is a group of order 3. Notice that we do not get back a group of the actual cosets, but instead we get a group *isomorphic* to the factor group.

Simple groups

It is easy to check to see if a group is void of any normal subgroups. The commands

```
sage: AlternatingGroup(5).is_simple()
True
sage: AlternatingGroup(4).is_simple()
False
```

prints True and then False.

Composition series

For any group, it is easy to obtain a composition series. There is an element of randomness in the algorithm, so you may not always get the same results. (But the list of factor groups is unique, according to the Jordan-Hölder theorem.) Also, the subgroups generated sometimes have more generators than necessary, so you might want to "study" each subgroup carefully by checking properties like its order.

An interesting example is:

```
DihedralGroup(105).composition_series()
```

The output will be a list of 5 subgroups of D_{105} , each a normal subgroup of its predecessor.

Several other series are possible, such as the derived series. Use tab-completion to see the possibilities.

Conjugacy

Given a group G, we can define a relation \sim on G by: for $a,b \in G$, $a \sim b$ if and only if there exists an element $g \in G$ such that $gag^{-1} = b$.

Since this is an equivalence relation, there is an associated partition of the elements of G into equivalence classes. For this very important relation, the classes are known as "conjugacy classes." A representative of each of these equivalence classes can be found as follows. Suppose G is a permutation group, then G.conjugacy_classes_representatives() will return a list of elements of G, one per conjugacy class.

Given an element $g \in G$, the "centralizer" of g is the set $C(g) = \{h \in G \mid hgh^{-1} = g\}$, which is a subgroup of G. A theorem tells us that the size of each conjugacy class is the order of the group divided by the order of the centralizer of an element of the class. With the following code we can determine the size of the conjugacy classes of the full symmetric group on 5 symbols:

```
sage: G = SymmetricGroup(5)
sage: group_order = G.order()
sage: reps = G.conjugacy_classes_representatives()
sage: class_sizes = []
sage: for g in reps:
....: class_sizes.append(group_order / G.centralizer(g).order())
...
sage: class_sizes
[1, 10, 15, 20, 20, 30, 24]
```

This should produce the list [1, 10, 15, 20, 20, 30, 24] which you can check sums to 120, the order of the group. You might be able to produce this list by counting elements of the group S_5 with identical cycle structure (which will require a few simple combinatorial arguments).

Sylow subgroups

Sylow's Theorems assert the existence of certain subgroups. For example, if p is a prime, and p^r divides the order of a group G, then G must have a subgroup of order p^r . Such a subgroup could be found among the output of the conjugacy_classes_subgroups () command by checking the orders of the subgroups produced. The map () command is a quick way to do this. The symmetric group on 7 symbols, S_7 , has order 7! = 5040 and is divisible by $2^4 = 16$. Let's find one example of a subgroup of permutations on 4 symbols with order 16:

```
sage: G = SymmetricGroup(7)
sage: subgroups = G.conjugacy_classes_subgroups()
sage: map(order, subgroups)
[1, 2, 2, 2, 3, 3, 4, 4, 4, 4, 4, 4, 4, 5, 6, 6, 6, 6, 6, 6, 6, 6, 7, 8, 8, 8, 8, 8, 8, 8, 9, 10, 10]
```

The map (order, subgroups) command will apply the order () function to each of the subgroups in the list subgroups. The output is thus a large list of the orders of many subgroups (96 to be precise). If you count carefully, you will see that the 49-th subgroup has order 16. You can retrieve this group for further study by referencing it as subgroups [48] (remember that counting starts at zero).

If p^r is the highest power of p to divide the order of G, then a subgroup of order p^r is known as a "Sylow p-subgroup." Sylow's Theorems also say any two Sylow p-subgroups are conjugate, so the output of conjugacy_classes_subgroups () should only contain each Sylow p-subgroup once. But there is an easier way, sylow_subgroup (p) will return one. Notice that the argument of the command is just the prime \$p\$, not the full power p^r . Failure to use a prime will generate an informative error message.

Groups of small order as permutation groups

We list here constructions, as permutation groups, for all of the groups of order less than 16.

```
Size Construction Notes

1 SymmetricGroup(1) Trivial
```

```
2
      SymmetricGroup(2)
                                                   Also CyclicPermutationGroup(2)
3
      CyclicPermutationGroup(3)
                                                   Prime order
4
      CyclicPermutationGroup(4)
                                                   Cvclic
4
      KleinFourGroup()
                                                   Abelian, non-cyclic
5
      CyclicPermutationGroup(5)
                                                   Prime order
6
      CyclicPermutationGroup(6)
                                                   Cyclic
6
      SymmetricGroup(3)
                                                   Non-abelian, also DihedralGroup(3)
7
     CyclicPermutationGroup(7)
                                                   Prime order
8
      CyclicPermutationGroup(8)
                                                   Cyclic
8
      D1 = CyclicPermutationGroup(4)
      D2 = CyclicPermutationGroup(2)
      G = direct_product_permgroups([D1,D2])
                                                   Abelian, non-cyclic
      D1 = CyclicPermutationGroup(2)
      D2 = CyclicPermutationGroup(2)
      D3 = CyclicPermutationGroup(2)
      G = direct_product_permgroups([D1,D2,D3])
                                                   Abelian, non-cyclic
8
      DihedralGroup (4)
                                                   Non-abelian
8
      QuaternionGroup()
                                                   Quaternions, also DiCyclicGroup(2)
9
      CyclicPermutationGroup(9)
                                                   Cyclic
      D1 = CyclicPermutationGroup(3)
      D2 = CyclicPermutationGroup(3)
      G = direct_product_permgroups([D1,D2])
                                                   Abelian, non-cyclic
10
     CyclicPermutationGroup(10)
                                                   Cvclic
10
     DihedralGroup(5)
                                                   Non-abelian
11
     CyclicPermutationGroup(11)
                                                   Prime order
12
      CyclicPermutationGroup(12)
                                                   Cyclic
12
      D1 = CyclicPermutationGroup(6)
      D2 = CyclicPermutationGroup(2)
      G = direct_product_permgroups([D1,D2])
                                                   Abelian, non-cyclic
12
      DihedralGroup(6)
                                                   Non-abelian
12
      AlternatingGroup(4)
                                                   Non-abelian, symmetries of tetrahedron
12
     DiCyclicGroup(3)
                                                   Non-abelian
                                                   Also semi-direct product $Z_3 \rtimes Z_4$
13
     CyclicPermutationGroup(13)
                                                   Prime order
14
      CyclicPermutationGroup(14)
                                                   Cyclic
14
     DihedralGroup(7)
                                                   Non-abelian
1.5
      CyclicPermutationGroup(15)
                                                   Cyclic
```

Acknowledgements

The construction of Sage is the work of many people, and the group theory portion is made possible by the extensive work of the creators of GAP. However, we will single out three people from the Sage team to thank for major contributions toward bringing you the group theory portion of Sage: David Joyner, William Stein, and Robert Bradshaw. Thanks!

12.1.5 Lie Methods and Related Combinatorics in Sage

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These notes explain how to use the mathematical software Sage for Lie group computations. Sage also contains many combinatorial algorithms. We will cover only some of these.

The Scope of this Document

Lie groups and algebras

Sage can be used to do standard computations for Lie groups and Lie algebras. The following categories of representations are equivalent:

- Complex representations of a compact, semisimple simply connected Lie group G.
- Complex representations of its Lie algebra g. This is a real Lie algebra, so representations are not required to be complex linear maps.
- Complex representations of its complexified Lie algebra $\mathfrak{g}_{\mathbf{C}} = \mathbf{C} \otimes \mathfrak{g}$. This is a complex Lie algebra and representations are required to be complex linear transformations.
- The complex analytic representations of the semisimple simply-connected complex analytic group $G_{\mathbf{C}}$ having $\mathfrak{g}_{\mathbf{C}}$ as its Lie algebra.
- Modules of the universal enveloping algebra $U(\mathfrak{g}_{\mathbf{C}})$.
- Modules of the quantized enveloping algebra $U_q(\mathfrak{g}_{\mathbf{C}})$.

For example, we could take G = SU(n), $\mathfrak{g} = \mathfrak{sl}(n, \mathbf{R})$, $\mathfrak{g}_{\mathbf{C}} = \mathfrak{sl}(n, \mathbf{C})$ and $G_{\mathbf{C}} = SL(n, \mathbf{C})$. Because these categories are the same, their representations may be studied simultaneously. The above equivalences may be expanded to include reductive groups like U(n) and GL(n) with a bit of care.

Here are some typical problems that can be solved using Sage:

- Decompose a module in any one of these categories into irreducibles.
- Compute the Frobenius-Schur indicator of an irreducible module.
- Compute the tensor product of two modules.
- If H is a subgroup of G, study the restriction of modules for G to H. The solution to this problem is called a branching rule.
- Find the multiplicities of the weights of the representation.

In addition to its representations, which we may study as above, a Lie group has various related structures. These include:

- The Weyl Group W.
- The Weight Lattice.
- · The Root System
- · The Cartan Type.
- The Dynkin diagram.
- The extended Dynkin diagram.

Sage contains methods for working with these structures.

If there is something you need that is not implemented, getting it added to Sage will likely be possible. You may write your own algorithm for an unimplemented task, and if it is something others will be interested in, it is probably possible to get it added to Sage.

Combinatorics

Sage supports a great many related mathematical objects. Some of these properly belong to combinatorics. It is beyond the scope of these notes to cover all the combinatorics in Sage, but we will try to touch on those combinatorial methods which have some connection with Lie groups and representation theory. These include:

- The affine Weyl group, an infinite group containing W.
- Kashiwara crystals, which are combinatorial analogs of modules in the above categories.
- · Coxeter group methods applicable to Weyl groups and the affine Weyl group, such as Bruhat order.
- The Iwahori Hecke algebras, which are deformations of the group algebras of W and the affine Weyl group.
- Kazhdan-Lusztig polynomials.

Lie Group Basics

Goals of this section

Since we must be brief here, this is not really a place to learn about Lie groups or Lie algebras. Rather, the point of this section is to outline what you need to know to use Sage effectively for Lie computations, and to fix ideas and notations.

Semisimple and reductive groups

If $g \in GL(n, \mathbb{C})$, then g may be uniquely factored as g_1g_2 where g_1 and g_2 commute, with g_1 semisimple (diagonalizable) and g_2 unipotent (all its eigenvalues equal to 1). This follows from the Jordan canonical form. If $g = g_1$ then g is called *semisimple* and if $g = g_2$ then g is called *unipotent*.

We consider a Lie group G and a class of representations such that if an element $g \in G$ is unipotent (resp. semisimple) in one faithful representation from the class, then it is unipotent (resp. semisimple) in every faithful representation of the class. Thus the notion of being semisimple or unipotent is intrinsic. Examples:

- Compact Lie groups with continuous representations
- Complex analytic groups with analytic representations
- Algebraic groups over **R** with algebraic representations.

A subgroup of G is called *unipotent* if it is connected and all its elements are unipotent. It is called a *torus* if it is connected, abelian, and all its elements are semisimple. The group G is called *reductive* if it has no nontrivial normal unipotent subgroup. For example, $GL(2, \mathbb{C})$ is reductive, but its subgroup:

$$\left\{ \left(\begin{array}{cc} a & b \\ & d \end{array}\right) \right\}$$

is not since it has a normal unipotent subgroup

$$\left\{\left(\begin{array}{cc} 1 & b \\ & 1 \end{array}\right)\right\}.$$

A group has a unique largest normal unipotent subgroup, called the *unipotent radical*, so it is reductive if and only if the unipotent radical is trivial.

A Lie group is called *semisimple* it is reductive and furthermore has no nontrivial normal tori. For example $GL(2, \mathbf{C})$ is reductive but not semisimple because it has a normal torus:

$$\left\{ \left(\begin{array}{cc} a \\ & a \end{array}\right) \right\}.$$

The group $SL(2, \mathbb{C})$ is semisimple.

Fundamental group and center

If G is a semisimple Lie group then its center and fundamental group are finite abelian groups. The universal covering group \tilde{G} is therefore a finite extension with the same Lie algebra. Any representation of G may be reinterpreted as a representation of the simply connected \tilde{G} . Therefore we may as well consider representations of \tilde{G} , and restrict ourselves to the simply connected group.

Parabolic subgroups and Levi subgroups

Let G be a reductive complex analytic group. A maximal solvable subgroup of G is called a *Borel subgroup*. All Borel subgroups are conjugate. Any subgroup P containing a Borel subgroup is called a *parabolic subgroup*. We may write P as a semidirect product of its maximal normal unipotent subgroup or *unipotent radical* P and a reductive subgroup M, which is determined up to conjugacy. The subgroup M is called a *Levi subgroup*.

Example: Let $G = GL_n(\mathbf{C})$ and let r_1, \ldots, r_k be integers whose sum is n. Then we may consider matrices of the form:

$$\begin{pmatrix}
g_1 & * & \cdots & * \\
& g_2 & & * \\
& & \ddots & \\
& & & g_r
\end{pmatrix}$$

where $g_i \in GL(r_i, \mathbf{C})$. The unipotent radical consists of the subgroup in which all $g_i = I_{r_i}$. The Levi subgroup (determined up to conjugacy) is:

$$M = \left\{ \begin{pmatrix} g_1 & & & \\ & g_2 & & \\ & & \ddots & \\ & & & g_r \end{pmatrix} \right\},$$

and is isomorphic to $M = GL(r_1, \mathbf{C}) \times \cdots \times GL(r_k, \mathbf{C})$. Therefore M is a Levi subgroup.

The notion of a Levi subgroup can be extended to compact Lie groups. Thus $U(r_1) \times \cdots \times U(r_k)$ is a Levi subgroup of U(n). However parabolic subgroups do not exist for compact Lie groups.

Cartan types

Semisimple Lie groups are classified by their *Cartan types*. There are both reducible and irreducible Cartan types in Sage. Let us start with the irreducible types. Such a type is implemented in Sage as a pair ['X', r] where 'X' is one of A, B, C, D, E, F or G and r is a positive integer. If 'X' is 'D' then we must have r > 1 and if 'X' is one of the *exceptional types* 'E', 'F' or 'G' then r is limited to only a few possibilities. The exceptional types are:

A simply-connected semisimple group is a direct product of simple Lie groups, which are given by the following table. The integer r is called the rank, and is the dimension of the maximal torus.

Here are the Lie groups corresponding to the classical types:

compact group	complex analytic group	Cartan type
SU(r+1)	$SL(r+1, \mathbf{C})$	A_r
spin(2r+1)	$spin(2r+1, \mathbf{C})$	B_r
Sp(2r)	$Sp(2r, \mathbf{C})$	C_r
spin(2r)	$spin(2r, \mathbf{C})$	D_r

You may create these Cartan types and their Dynkin diagrams as follows:

```
sage: ct = CartanType("D5"); ct
['D', 5]
```

Here "D5" is an abbreviation for ['D', 5]. The group spin(n) is the simply-connected double cover of the orthogonal group SO(n).

Dual Cartan types

Every Cartan type has a dual, which you can get from within Sage:

```
sage: CartanType("B4").dual()
['C', 4]
```

Types other than B_r and C_r for r > 2 are self-dual in the sense that the dual is isomorphic to the original type; however the isomorphism of a Cartan type with its dual might relabel the vertices. We can see this as follows:

```
sage: CartanType("F4").dynkin_diagram()
O---O=>=O---O
1    2    3    4
F4
sage: CartanType("F4").dual()
['F', 4] relabelled by {1: 4, 2: 3, 3: 2, 4: 1}
sage: CartanType("F4").dual().dynkin_diagram()
O---O=>=O---O
4    3    2    1
F4 relabelled by {1: 4, 2: 3, 3: 2, 4: 1}
```

Reducible Cartan types

If G is a Lie group of finite index in $G_1 \times G_2$, where G_1 and G_2 are Lie groups of positive dimension, then G is called *reducible*. In this case, the root system of G is the disjoint union of the root systems of G_1 and G_2 , which lie in orthogonal subspaces of the ambient space of the weight space of G. The Cartan type of G is thus *reducible*.

Reducible Cartan types are supported in Sage as follows:

```
sage: RootSystem("A1xA1")
Root system of type A1xA1
sage: WeylCharacterRing("A1xA1")
The Weyl Character Ring of Type A1xA1 with Integer Ring coefficients
```

Low dimensional Cartan types

There are some isomorphisms that occur in low degree.

Cartan Type	Group	Equivalent Type	Isomorphic Group
B_2	spin(5)	C_2	Sp(4)
D_3	spin(6)	A_3	SL(4)
D_2	spin(4)	$A1 \times A_1$	$SL(2) \times SL(2)$
B_1	spin(3)	A_1	SL(2)
C_1	Sp(2)	A_1	SL(2)

Sometimes the redundant Cartan types such as D_3 and D_2 are excluded from the list of Cartan types. However Sage allows them since excluding them leads to exceptions having to be made in algorithms. A better approach, which is followed by Sage, is to allow the redundant Cartan types, but to implement the isomorphisms explicitly as special cases of branching rules. The utility of this approach may be seen by considering that the rank one group SL(2) has different natural weight lattices realizations depending on whether we consider it to be SL(2), Spin(2) or Sp(2):

```
sage: RootSystem("A1").ambient_space().simple_roots()
Finite family {1: (1, -1)}
sage: RootSystem("B1").ambient_space().simple_roots()
Finite family {1: (1)}
sage: RootSystem("C1").ambient_space().simple_roots()
Finite family {1: (2)}
```

Relabeled Cartan types

By default Sage uses the labeling of the Dynkin diagram from [Bourbaki46]. There is another labeling of the vertices due to Dynkin. Most of the literature follows [Bourbaki46], though [Kac] follows Dynkin.

If you need to use Dynkin's labeling, you should be aware that Sage does support relabeled Cartan types. See the documentation in sage.combinat.root_system.type_relabel for further information.

Standard realizations of the ambient spaces

These realizations follow the Appendix in [Bourbaki46]. See the *Root system plot tutorial* for how to visualize them.

Type A For type A_r we use an r+1 dimensional ambient space. This means that we are modeling the Lie group U(r+1) or $GL(r+1, \mathbb{C})$ rather than SU(r+1) or $SL(r+1, \mathbb{C})$. The ambient space is identified with \mathbb{Q}^{r+1} :

```
sage: RootSystem("A3").ambient_space().simple_roots()
Finite family {1: (1, -1, 0, 0), 2: (0, 1, -1, 0), 3: (0, 0, 1, -1)}
sage: RootSystem("A3").ambient_space().fundamental_weights()
Finite family {1: (1, 0, 0, 0), 2: (1, 1, 0, 0), 3: (1, 1, 1, 0)}
sage: RootSystem("A3").ambient_space().rho()
(3, 2, 1, 0)
```

The dominant weights consist of integer r+1-tuples $\lambda=(\lambda_1,\ldots,\lambda_{r+1})$ such that $\lambda_1\geq\cdots\geq\lambda_{r+1}$.

See *SL versus GL* for further remarks about Type A.

Type B For the remaining classical Cartan types B_r , C_r and D_r we use an r-dimensional ambient space:

```
sage: RootSystem("B3").ambient_space().simple_roots()
Finite family {1: (1, -1, 0), 2: (0, 1, -1), 3: (0, 0, 1)}
sage: RootSystem("B3").ambient_space().fundamental_weights()
Finite family {1: (1, 0, 0), 2: (1, 1, 0), 3: (1/2, 1/2, 1/2)}
sage: RootSystem("B3").ambient_space().rho()
(5/2, 3/2, 1/2)
```

This is the Cartan type of spin(2r+1). The last fundamental weight (1/2, 1/2, ..., 1/2) is the highest weight of the 2^r dimensional *spin representation*. All the other fundamental representations factor through the homomorphism $spin(2r+1) \rightarrow SO(2r+1)$ and are representations of the orthogonal group.

The dominant weights consist of r-tuples of integers or half-integers $(\lambda_1, \dots, \lambda_r)$ such that $\lambda_1 \geq \lambda_2 \dots \geq \lambda_r \geq 0$, and such that the differences $\lambda_i - \lambda_j \in \mathbf{Z}$.

Type C

```
sage: RootSystem("C3").ambient_space().simple_roots()
Finite family {1: (1, -1, 0), 2: (0, 1, -1), 3: (0, 0, 2)}
sage: RootSystem("C3").ambient_space().fundamental_weights()
Finite family {1: (1, 0, 0), 2: (1, 1, 0), 3: (1, 1, 1)}
sage: RootSystem("C3").ambient_space().rho()
(3, 2, 1)
```

This is the Cartan type of the symplectic group Sp(2r).

The dominant weights consist of r-tuples of integers $\lambda = (\lambda_1, \dots, \lambda_{r+1})$ such that $\lambda_1 \ge \dots \ge \lambda_r \ge 0$.

Type D

```
sage: RootSystem("D4").ambient_space().simple_roots()
Finite family {1: (1, -1, 0, 0), 2: (0, 1, -1, 0), 3: (0, 0, 1, -1), 4: (0, 0, 1, 1)}
sage: RootSystem("D4").ambient_space().fundamental_weights()
Finite family {1: (1, 0, 0, 0), 2: (1, 1, 0, 0), 3: (1/2, 1/2, 1/2, -1/2), 4: (1/2, 1/2, 1/2, 1/2)}
sage: RootSystem("D4").ambient_space().rho()
(3, 2, 1, 0)
```

This is the Cartan type of spin(2r). The last two fundamental weights are the highest weights of the two 2^{r-1} -dimensional spin representations.

The dominant weights consist of r-tuples of integers $\lambda = (\lambda_1, \dots, \lambda_{r+1})$ such that $\lambda_1 \ge \dots \ge \lambda_{r-1} \ge |\lambda_r|$.

Exceptional Types We leave the reader to examine the exceptional types. You can use Sage to list the fundamental dominant weights and simple roots.

Weights and the ambient space

Let G be a reductive complex analytic group. Let T be a maximal torus, $\Lambda = X^*(T)$ be its group of analytic characters. Then $T \cong (\mathbf{C}^{\times})^r$ for some r and $\Lambda \cong \mathbf{Z}^r$.

Example 1: Let $G = \operatorname{GL}_{r+1}(\mathbf{C})$. Then T is the diagonal subgroup and $X^*(T) \cong \mathbf{Z}^{r+1}$. If $\lambda = (\lambda_1, \dots, \lambda_n)$ then λ is identified with the rational character

$$\mathbf{t} = \left(egin{array}{ccc} t_1 & & & \ & \ddots & & \ & & t_n \end{array}
ight) \longmapsto \prod t_i^{\lambda_i}.$$

Example 2: Let $G = \operatorname{SL}_{r+1}(\mathbf{C})$. Again T is the diagonal subgroup but now if $\lambda \in \mathbf{Z}^{\Delta} = \{(d, \dots, d) | d \in \mathbf{Z}\} \subseteq \mathbf{Z}^{r+1}$ then $\prod t_i^{\lambda_i} = \det(\mathbf{t})^d = 1$, so $X^*(T) \cong \mathbf{Z}^{r+1}/\mathbf{Z}^{\Delta} \cong \mathbf{Z}^r$.

- Elements of Λ are called *weights*.
- If $\pi:G\to GL(V)$ is any representation we may restrict π to T. Then the characters of T that occur in this restriction are called the *weights of* π .

- G acts on its Lie algebra by conjugation (the *adjoint representation*).
- The nonzero weights of the adjoint representation are called *roots*.
- The *ambient space* of Λ is $\mathbf{Q} \otimes \Lambda$.

The root system

As we have mentioned, G acts on its complexified Lie algebra $\mathfrak{g}_{\mathbf{C}}$ by the adjoint representation. The zero weight space $\mathfrak{g}_{\mathbf{C}}(0)$ is just the Lie algebra of T itself. The other nonzero weights each appear with multiplicity one and form an interesting configuration of vectors called the *root system* Φ .

It is convenient to partition Φ into two sets Φ^+ and Φ^- such that Φ^+ consists of all roots lying on one side of a hyperplane. Often we arrange things so that G is embedded in $GL(n, \mathbf{C})$ in such a way that the positive weights correspond to upper triangular matrices. Thus if α is a positive root, its weight space $\mathfrak{g}_{\mathbf{C}}(\alpha)$ is spanned by a vector X_{α} , and the exponential of this eigenspace in G is a one-parameter subgroup of unipotent matrices. It is always possible to arrange that this one-parameter subgroup consists of upper triangular matrices.

If α is a positive root that cannot be decomposed as a sum of other positive roots, then α is called a *simple root*. If G is semisimple of rank r, then r is the number of positive roots. Let $\alpha_1, \ldots, \alpha_r$ be these.

The Weyl group

Let G be a complex analytic group. Let T be a maximal torus, and let N(T) be its normalizer. Let W = N(T)/T be the Weyl group. It acts on T by conjugation; therefore it acts on the weight lattice Λ and its ambient space. The ambient space admits an inner product that is invariant under this action. Let (v|w) denote this inner product. If α is a root let r_{α} denote the reflection in the hyperplane of the ambient space that is perpendicular to α . If $\alpha = \alpha_i$ is a simple root, then we use the notation s_i to denote r_{α} .

Then s_1, \ldots, s_r generate W, which is a *Coxeter group*. This means that it is generated by elements s_i of order two and that if m(i, j) is the order of $s_i s_j$, then

$$W = \left\langle s_i \mid s_i^2 = 1, (s_i s_j)^{m(i,j)} = 1 \right\rangle$$

is a presentation. An important function $\ell:W\to \mathbf{Z}$ is the *length* function, where $\ell(w)$ is the length of the shortest decomposition of w into a product of simple reflections.

The dual root system

The *coroots* are certain linear functionals on the ambient space that also form a root system. Since the ambient space admits a W-invariant inner product $(\ |\)$, they may be identified with elements of the ambient space itself. Then they are proportional to the roots, though if the roots have different lengths, long roots correspond to short coroots and conversely. The coroot corresponding to the root α is

$$\alpha^{\vee} = \frac{2\alpha}{(\alpha|\alpha)}.$$

We can also describe the natural pairing between coroots and roots using this invariant inner product as

$$\langle \alpha^{\vee}, \beta \rangle = 2 \frac{(\alpha | \beta)}{(\alpha | \alpha)}.$$

The Dynkin diagram

The Dynkin diagram is a graph whose vertices are in bijection with the set simple roots. We connect the vertices corresponding to roots that are not orthogonal. Usually two such roots (vertices) make an angle of $2\pi/3$, in which case we connect them with a single bond. Occasionally they may make an angle of $3\pi/4$ in which case we connect them with a double bond, or $5\pi/6$ in which case we connect them with a triple bond. If the bond is single, the roots have the same length with respect to the inner product on the ambient space. In the case of a double or triple bond, the two simple roots in questions have different length, and the bond is drawn as an arrow from the long root to the short root. Only the exceptional group G_2 has a triple bond.

There are various ways to get the Dynkin diagram in Sage:

```
sage: DynkinDiagram("D5")
       0 5
       0---0
1
   2 3 4
sage: ct = CartanType("E6"); ct
['E', 6]
sage: ct.dynkin_diagram()
   -0---0---0
      4 5
   3
1
sage: B4 = WeylCharacterRing("B4"); B4
The Weyl Character Ring of Type B4 with Integer Ring coefficients
sage: B4.dynkin_diagram()
0---0-->=0
   2 3 4
1
В4
sage: RootSystem("G2").dynkin_diagram()
 3
0=<=0
1
   2
G2
```

The Cartan matrix

Consider the natural pairing \langle , \rangle between coroots and roots, then the defining matrix of this pairing is called the *Cartan matrix*. That is to say, the Cartan matrix $A = (a_{ij})_{ij}$ is given by

$$a_{ij} = \langle \alpha_i^{\vee}, \alpha_j \rangle.$$

This uniquely corresponds to a root system/Dynkin diagram/Lie group.

We note that we have made a convention choice, and the opposite convention corresponds to taking the transpose of the Cartan matrix.

Fundamental weights and the Weyl vector

There are certain weights $\omega_1, \ldots, \omega_r$ that:

$$\langle \omega_j, \alpha_i \rangle = 2 \frac{(\alpha_i | \omega_j)}{(\alpha_i | \alpha_i)} = \delta_{ij}.$$

If G is semisimple then these are uniquely determined, whereas if G is reductive but not semisimple we may choose them conveniently.

Let ρ be the sum of the fundamental dominant weights. If G is semisimple, then ρ is half the sum of the positive roots. In case G is not semisimple, we have noted, the fundamental weights are not completely determined by the inner product condition given above. If we make a different choice, then ρ is altered by a vector that is orthogonal to all roots. This is a harmless change for many purposes such as the Weyl character formula.

In Sage, this issue arises only for Cartan type A_r . See SL versus GL.

Representations and characters

Let T be a maximal torus and $\Lambda = X^*(T)$ be the group of rational characters. Then $\Lambda \cong \mathbf{Z}^r$.

- Recall that elements of $\Lambda \cong \mathbf{Z}^r$ are called *weights*.
- The Weyl group W = N(T)/T acts on T, hence on Λ and its ambient space by conjugation.
- The ambient space $\mathbf{Q} \otimes X^*(T) \cong \mathbf{Q}^r$ has a fundamental domain \mathcal{C}^+ for the Weyl group W called the *positive Weyl chamber*. Weights in \mathcal{C}^+ are called *dominant*.
- Then C^+ consists of all vectors such that $(\alpha|v) \geq 0$ for all positive roots α .
- It is useful to embed Λ in \mathbb{R}^r and consider weights as lattice points.
- If (π, V) is a representation then restricting to T, the module V decomposes into a direct sum of weight eigenspaces $V(\mu)$ with multiplicity $m(\mu)$ for weight μ .
- There is a unique highest weight λ with respect to the partial order. We have $\lambda \in \mathcal{C}$ and $m(\lambda) = 1$.
- $V \longleftrightarrow \lambda$ gives a bijection between irreducible representations and weights λ in \mathcal{C}^+ .

Assuming that G is simply-connected (or more generally, reductive with a simply-connected derived group) every dominant weight λ is the highest weight of a unique irreducible representation π_{λ} , and $\lambda \mapsto \pi_{\lambda}$ gives a parametrization of the isomorphism classes of irreducible representations of G by the dominant weights.

The *character* of π_{λ} is the function $\chi_{\lambda}(g) = tr(\pi_{\lambda}(g))$. It is determined by its values on T. If $(z) \in T$ and $\mu \in \Lambda$, let us write \mathbf{z}^{μ} for the value of μ on \mathbf{z} . Then the character:

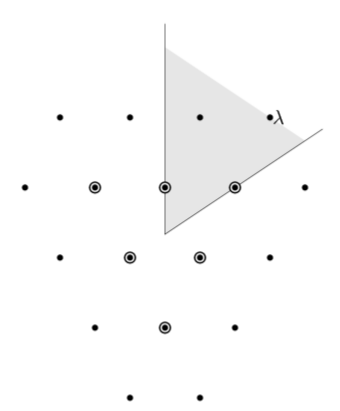
$$\chi_{\lambda}(\mathbf{z}) = \sum_{\mu \in \Lambda} m(\mu) \, \mathbf{z}^{\lambda}.$$

Sometimes this is written

$$\chi_{\lambda} = \sum_{\mu \in \Lambda} m(\mu) \, e^{\lambda}.$$

The meaning of e^{λ} is subject to interpretation, but we may regard it as the image of the additive group Λ in its group algebra. The character is then regarded as an element of this ring, the group algebra of Λ .

Representations: an example



In this example, $G = \mathrm{SL}(3, \mathbf{C})$. We have drawn the weights of an irreducible representation with highest weight λ . The shaded region is \mathcal{C}^+ . λ is a dominant weight, and the labeled vertices are the weights with positive multiplicity in $V(\lambda)$. The weights weights on the outside have $m(\mu) = 1$, while the six interior weights (with double circles) have $m(\mu) = 2$.

Partitions and Schur polynomials

The considerations of this section are particular to type A. We review the relationship between characters of $GL(n, \mathbf{C})$ and symmetric function theory.

A partition λ is a sequence of descending nonnegative integers:

$$\lambda = (\lambda_1, \lambda_2, \dots, \lambda_n), \quad \lambda_1 \ge \lambda_2 \ge \dots \ge \lambda_n \ge 0.$$

We do not distinguish between two partitions if they differ only by some trailing zeros, so (3,2)=(3,2,0). If l is the last integer such that $\lambda_l>0$ then we say that l is the *length* of λ . If $k=\sum \lambda_i$ then we say that λ is a *partition* of k and write $\lambda \vdash k$.

A partition of length $\leq n = r+1$ is therefore a dominant weight of type $\lceil ' A', r \rceil$. Not every dominant weight is a partition, since the coefficients in a dominant weight could be negative. Let us say that an element $\mu = (\mu_1, \mu_2, \cdots, \mu_n)$ of the $\lceil ' A', r \rceil$ root lattice is *effective* if the $\mu_i \geq 0$. Thus an effective dominant weight of $\lceil ' A', r \rceil$ is a partition of length $\leq n$, where n = r + 1.

Let λ be a dominant weight, and let χ_{λ} be the character of $GL(n, \mathbf{C})$ with highest weight λ . If k is any integer we may consider the weight $\mu = (\lambda_1 + k, \dots, \lambda_n + k)$ obtained by adding k to each entry. Then $\chi_{\mu} = \det^k \otimes \chi_{\lambda}$. Clearly by choosing k large enough, we may make μ effective.

So the characters of irreducible representations of $GL(n, \mathbf{C})$ do not all correspond to partitions, but the characters indexed by partitions (effective dominant weights) are enough that we can write any character $\det^{-k} \chi_{\mu}$ where μ is a

partition. If we take $k = -\lambda_n$ we could also arrange that the last entry in λ is zero.

If λ is an effective dominant weight, then every weight that appears in χ_{λ} is effective. (Indeed, it lies in the convex hull of $w(\lambda)$ where w runs through the Weyl group $W = S_n$.) This means that if

$$g = \begin{pmatrix} z_1 & & \\ & \ddots & \\ & & z_n \end{pmatrix} \in GL(n, \mathbf{C})$$

then $\chi_{\lambda}(g)$ is a polynomial in the eigenvalues of g. This is the Schur polynomial $s_{\lambda}(z_1,\ldots,z_n)$.

Affine Cartan types

There are also affine Cartan types, which correspond to (infinite dimensional) affine Lie algebras. There are affine Cartan types of the form ['X', r, 1] if X=A,B,C,D,E,F,G and ['X', r] is an ordinary Cartan type. There are also *twisted affine types* of the form [X, r, k], where k=2 or 3 if the Dynkin diagram of the ordinary Cartan type [X, r] has an automorphism of degree k. When k=1, the affine Cartan type is said to be *untwisted*.

Illustrating some of the methods available for the untwisted affine Cartan type ['A', 4, 1]:

The twisted affine Cartan types are relabeling of the duals of certain untwisted Cartan types:

```
sage: CartanType(['A',3,2])
['B', 2, 1]^*
sage: CartanType(['D',4,3])
['G', 2, 1]^* relabelled by {0: 0, 1: 2, 2: 1}
```

The affine root and the extended Dynkin diagram

For the extended Dynkin diagram, we add one negative root α_0 . For the untwisted types, this is the root whose negative is the highest weight in the adjoint representation. Sometimes this is called the *affine root*. We make the Dynkin diagram as before by measuring the angles between the roots. This extended Dynkin diagram is useful for many purposes, such as finding maximal subgroups and for describing the affine Weyl group.

In particular, the hyperplane for the reflection r_0 , used in generating the affine Weyl group is translated off the origin (so it becomes an affine hyperplane). Now the root system is not described as linear transformations on an Euclidean space, but instead by *affine* transformations. Thus the dominant chamber has finite volume and tiles the Eucledian space. Moreover, each such tile corresponds to a unique element in the affine Weyl group.

The extended Dynkin diagram may be obtained as the Dynkin diagram of the corresponding untwisted affine type:

The extended Dynkin diagram is also a method of the WeylCharacterRing:

We note the following important distinctions from the classical cases:

- The affine Weyl groups are all infinte.
- Type $A_1^{(1)}$ has two anti-parallel roots with distinct reflections. The Dynkin diagram in this case is represented by a double bond with arrows going in both directions.

Twisted affine root systems

For the construction of α_0 in the twisted types, we refer the reader to Chaper 8 of [Kac]. As mentioned above, most twisted types can be constructed by the taking the dual root system of an untwisted type. However the type $A_{2n}^{(2)}$ root system which can only be constructed by the twisting procedure defined in [Kac]. It has the following properties:

- The Dynkin diagram of type $A_2^{(2)}$ has a quadruple bond with an arrow pointing from the short root to the long root.
- Type $A_{2n}^{(2)}$ for n>1 has 3 different root lengths.

Further Generalizations

If a root system (on an Euclidean space) has only the angles $\pi/2$, $2\pi/3$, $3\pi/4$, $5\pi/6$ between its roots, then we call the root system *crystallographic* (on Wikipedia article Root_system, this condition is called integrality since for any two roots we have $\langle \beta, \alpha \rangle \in \mathbf{Z}$). So if we look at the reflection group generated by the roots (this is not a Weyl group), we get general Coxeter groups (with non-infinite labels) and non-crystallographic Coxeter groups are not connected with Lie theory.

However we can generalize Dynkin diagrams (equivalently Cartan matrices) to have all its edges labelled by (a,b) where $a,b \in \mathbf{Z}_{>0}$ and corresponds to having a arrows point one way and b arrows pointing the other. For example in type $A_1^{(1)}$, we have one edge of (2,2), or in type $A_2^{(2)}$, we have one edge of (1,4) (equivalently (4,1)). These edge

label between i and j corresponds to the entries a_{ij} and a_{ji} in the Cartan matrix. These are used to construct a class of (generally infinite dimensional) Lie algebras called Kac-Moody (Lie) algebras, which in turn are used to construct quantum groups. We refer the reader to [Kac] and [HongKang2002] for more information.

The Weyl Character Ring

Weyl character rings

The Weyl character ring is the representation ring of a compact Lie group. It has a basis consisting of the irreducible representations of G, or equivalently, their characters. The addition and multiplication in the Weyl character ring correspond to direct sum and tensor product of representations.

Methods of the ambient space

In Sage, many useful features of the Lie group are available as methods of the ambient space:

```
sage: S = RootSystem("B2").ambient_space(); S
Ambient space of the Root system of type ['B', 2]
sage: S.roots()
[(1, -1), (1, 1), (1, 0), (0, 1), (-1, 1), (-1, -1), (-1, 0), (0, -1)]
sage: S.fundamental_weights()
Finite family {1: (1, 0), 2: (1/2, 1/2)}
sage: S.positive_roots()
[(1, -1), (1, 1), (1, 0), (0, 1)]
sage: S.weyl_group()
Weyl Group of type ['B', 2] (as a matrix group acting on the ambient space)
```

Methods of the Weyl character ring

If you are going to work with representations, you may want to create a Weyl Character ring. Many methods of the ambient space are available as methods of the Weyl character ring:

Other useful methods of the Weyl character ring include:

- cartan_type
- highest_root
- positive root
- extended_dynkin_diagram
- rank

Some methods of the ambient space are not implemented as methods of the Weyl character ring. However, the ambient space itself is a method, and so you have access to its methods from the Weyl character ring:

```
sage: B3 = WeylCharacterRing("B3")
sage: B3.space().weyl_group()
Weyl Group of type ['B', 3] (as a matrix group acting on the ambient space)
sage: B3.space()
Ambient space of the Root system of type ['B', 3]
sage: B3.space().rho()
(5/2, 3/2, 1/2)
sage: B3.cartan_type()
['B', 3]
```

Coroot notation

It is useful to give the Weyl character ring a name that corresponds to its Cartan type. This has the effect that the ring can parse its own output:

```
sage: G2 = WeylCharacterRing("G2")
sage: [G2(fw) for fw in G2.fundamental_weights()]
[G2(1,0,-1), G2(2,-1,-1)]
sage: G2(1,0,-1)
G2(1,0,-1)
```

Actually the prefix for the ring is configurable, so you don't really have to call this ring G2. Type WeylCharacterRing? at the sage: prompt for details.

There is one important option that you may want to know about. This is *coroot notation*. You select this by specifying the option style="coroots" when you create the ring. With the coroot style, the fundamental weights are represented $(1,0,0,\ldots)$, $(0,1,0,\ldots)$ instead of as vectors in the ambient space:

```
sage: B3 = WeylCharacterRing("B3", style="coroots")
sage: [B3(fw) for fw in B3.fundamental_weights()]
[B3(1,0,0), B3(0,1,0), B3(0,0,1)]
sage: B3(0,0,1)
B3(0,0,1)
sage: B3(0,0,1).degree()
```

The last representation is the eight dimensional spin representation of G = spin(7), the double cover of the orthogonal group SO(7). In the default notation it would be represented B3 (1/2, 1/2, 1/2).

With the coroot notation every irreducible representation is represented B3 (a, b, c) where a, b and c are nonnegative integers. This is often convenient. For many purposes the coroot style is preferable.

One disadvantage: in the coroot style the Lie group or Lie algebra is treated as semisimple, so you lose the distinction between GL(n) and SL(n); you also some information about representations of E6 and E7 for the same reason.

Tensor products of representations

The multiplication in the Weyl character ring corresponds to tensor product. This gives us a convenient way of decomposing a tensor product into irreducibles:

```
sage: B3 = WeylCharacterRing("B3")
sage: fw = B3.fundamental_weights()
sage: spinweight = fw[3]; spinweight
```

```
(1/2, 1/2, 1/2)
sage: spin = B3(spinweight); spin
B3(1/2,1/2,1/2)
sage: spin.degree()
```

The element spin of the WeylCharacterRing is the representation corresponding to the third highest weight representation, the eight-dimensional spin representation of spin(7). We could just as easily construct it with the command:

```
sage: spin = B3(1/2, 1/2, 1/2)
```

We may compute its tensor product with itself, using the multiplicative structure of the Weyl character ring:

```
sage: chi = spin*spin; chi
B3(0,0,0) + B3(1,0,0) + B3(1,1,0) + B3(1,1,1)
```

We have taken the eight-dimensional spin representation and tensored with itself. We see that the tensor square splits into four irreducibles, each with multiplicity one.

The highest weights that appear here are available (with their coefficients) through the usual free module accessors:

```
sage: from pprint import pprint
sage: list(chi)
                                           # random
[((1, 1, 1), 1), ((0, 0, 0), 1), ((1, 0, 0), 1), ((1, 1, 0), 1)]
sage: sorted(chi, key=str)
[((0, 0, 0), 1), ((1, 0, 0), 1), ((1, 1, 0), 1), ((1, 1, 1), 1)]
sage: pprint(dict(chi))
\{(0, 0, 0): 1, (1, 0, 0): 1, (1, 1, 0): 1, (1, 1, 1): 1\}
sage: M = sorted(chi.monomials(), key=lambda x: x.support()); M
[B3(0,0,0), B3(1,0,0), B3(1,1,0), B3(1,1,1)]
sage: sorted(chi.support())
[(0, 0, 0), (1, 0, 0), (1, 1, 0), (1, 1, 1)]
sage: chi.coefficients()
[1, 1, 1, 1]
sage: [r.degree() for r in M]
[1, 7, 21, 35]
sage: sum(r.degree() for r in chi.monomials())
64
```

Here we have extracted the individual representations, computed their degrees and checked that they sum up to 64.

Weight multiplicities

The weights of the character are available (with their coefficients) through the method weight_multiplicities. Continuing from the example in the last section:

```
sage: pprint(chi.weight_multiplicities())
{(0, 0, 0): 8, (-1, 0, 0): 4, (-1, -1, 0): 2, (-1, -1, -1): 1,
    (-1, -1, 1): 1, (-1, 1, 0): 2, (-1, 1, -1): 1, (-1, 1, 1): 1,
    (-1, 0, -1): 2, (-1, 0, 1): 2, (1, 0, 0): 4, (1, -1, 0): 2,
    (1, -1, -1): 1, (1, -1, 1): 1, (1, 1, 0): 2, (1, 1, -1): 1,
    (1, 1, 1): 1, (1, 0, -1): 2, (1, 0, 1): 2, (0, -1, 0): 4,
    (0, -1, -1): 2, (0, -1, 1): 2, (0, 1, 0): 4, (0, 1, -1): 2,
    (0, 1, 1): 2, (0, 0, -1): 4, (0, 0, 1): 4}
```

Each key of this dictionary is a weight, and its value is the multiplicity of that weight in the character.

Example

Suppose that we wish to compute the integral

$$\int_{U(n)} |tr(g)|^{2k} \, dg$$

for various n. Here U(n) is the unitary group, which is the maximal compact subroup of $GL(n, \mathbb{C})$. The irreducible unitary representations of U(n) may be regarded as the basis elements of the WeylCharacterRing of type A_r , where r=n-1 so we might work in that ring. The trace tr(g) is then just the character of the standard representation. We may realize it in the WeylCharacterRing by taking the first fundamental weight and coercing it into the ring. For example, if k=5 and n=3 so r=2:

```
sage: A2 = WeylCharacterRing("A2")
sage: fw = A2.fundamental_weights(); fw
Finite family {1: (1, 0, 0), 2: (1, 1, 0)}
sage: tr = A2(fw[1]); tr
A2(1,0,0)
```

We may compute the norm square the character tr^5 by decomposing it into irreducibles, and taking the sum of the squares of their multiplicities. By Schur orthogonality, this gives the inner product of the $tr(g)^5$ with itself, that is, the integral of $|tr(g)|^{10}$:

```
sage: sum(d^2 for d in (tr^5).coefficients())
103
```

So far we have been working with n = 3. For general n:

```
sage: def f(n,k):
....: R = WeylCharacterRing(['A',n-1])
....: tr = R(R.fundamental_weights()[1])
....: return sum(d^2 for d in (tr^k).coefficients())
sage: [f(n,5) for n in [2..7]]
[42, 103, 119, 120, 120, 120]
```

We see that the 10-th moment of tr(g) is just 5! when n is sufficiently large. What if we fix n and vary k?

```
sage: [f(2,k) for k in [1..10]]
[1, 2, 5, 14, 42, 132, 429, 1430, 4862, 16796]
sage: [catalan_number(k) for k in [1..10]]
[1, 2, 5, 14, 42, 132, 429, 1430, 4862, 16796]
```

Frobenius-Schur indicator

The Frobeinus-Schur indicator of an irreducible representation of a compact Lie group G with character χ is:

$$\int_G \chi(g^2) \, dg$$

The Haar measure is normalized so that vol(G) = 1. The Frobenius-Schur indicator equals 1 if the representation is real (orthogonal), -1 if it is quaternionic (symplectic) and 0 if it is complex (not self-contragredient). This is a method of weight ring elements corresponding to irreducible representations. Let us compute the Frobenius-Schur indicators of the spin representations of some odd spin groups:

```
sage: def spinrepn(r):
....: R = WeylCharacterRing(['B',r])
....: return R(R.fundamental_weights()[r])
....:
sage: spinrepn(3)
B3(1/2,1/2,1/2)
sage: for r in [1..4]: print r, spinrepn(r).frobenius_schur_indicator()
1 -1
2 -1
3 1
4 1
```

Here we have defined a function that returns the spin representation of the group spin(2r+1) with Cartan type [B', r], then computed the Frobenius-Schur indicators for a few values. From this experiment we see that the spin representations of spin(3) and spin(5) are symplectic, while those of spin(7) and spin(9) are orthogonal.

Symmetric and exterior powers

Sage can compute symmetric and exterior powers of a representation:

```
sage: B3 = WeylCharacterRing("B3", style="coroots")
sage: spin = B3(0,0,1); spin.degree()
8
sage: spin.exterior_power(2)
B3(1,0,0) + B3(0,1,0)
sage: spin.exterior_square()
B3(1,0,0) + B3(0,1,0)
sage: spin.exterior_power(5)
B3(0,0,1) + B3(1,0,1)
sage: spin.symmetric_power(5)
B3(0,0,1) + B3(0,0,3) + B3(0,0,5)
```

The k-th exterior square of a representation is zero if k is greater than the degree of the representation. However the k-th symmetric power is nonzero for all k.

The tensor square of any representation decomposes as the direct sum of the symmetric and exterior squares:

```
sage: C4 = WeylCharacterRing("C4",style="coroots")
sage: chi = C4(1,0,0,0); chi.degree()
8
sage: chi.symmetric_square()
C4(2,0,0,0)
sage: chi.exterior_square()
C4(0,0,0,0) + C4(0,1,0,0)
sage: chi^2 == chi.symmetric_square() + chi.exterior_square()
True
```

Since in this example the exterior square contains the trivial representation we expect the Frobenius-Schur indicator to be -1, and indeed it is:

```
sage: chi = C4(1,0,0,0)
sage: chi.frobenius_schur_indicator()
-1
```

This is not surprising since this is the standard representation of a symplectic group, which is symplectic by definition!

Weyl dimension formula

If the representation is truly large you will not be able to construct it in the Weyl character ring, since internally it is represented by a dictionary of its weights. If you want to know its degree, you can still compute that since Sage implements the Weyl dimension formula. The degree of the representation is implemented as a method of its highest weight vector:

```
sage: L = RootSystem("E8").ambient_space()
sage: [L.weyl_dimension(f) for f in L.fundamental_weights()]
[3875, 147250, 6696000, 6899079264, 146325270, 2450240, 30380, 248]
```

It is a fact that for any compact Lie group if ρ is the Weyl vector (half the sum of the positive roots) then the degree of the irreducible representation with highest weight ρ equals 2^N where N is the number of positive roots. Let us check this for E_8 . In this case N=120:

```
sage: L = RootSystem("E8").ambient_space()
sage: len(L.positive_roots())
120
sage: 2^120
1329227995784915872903807060280344576
sage: L.weyl_dimension(L.rho())
1329227995784915872903807060280344576
```

SL versus GL

Sage takes the weight space for type ['A', r] to be r + 1 dimensional. As a biproduct, if you create the Weyl character ring with the command:

```
sage: A2 = WeylCharacterRing("A2")
```

Then you are effectively working with GL(3) instead of SL(3). For example, the determinant is the character A2 (1,1,1). However, as we will explain later, you can work with SL(3) if you like, so long as you are willing to work with fractional weights. On the other hand if you create the Weyl character ring with the command:

```
sage: A2 = WeylCharacterRing("A2", style="coroots")
```

Then you are working with SL(3).

There are some advantages to this arrangement:

- The group GL(r+1) arises frequently in practice. For example, even if you care mainly about semisimple groups, the group GL(r+1) may arise as a Levi subgroup.
- It avoids fractional weights. If you want to work with SL(3) the fundamental weights are (2/3, -1/3, -1/3) and (1/3, 1/3, -2/3). If you work instead with GL(3), they are (1, 0, 0) and (1, 1, 0). For many mathematical purposes it doesn't make any difference which you use. This is because the difference between (2/3, -1/3, -1/3) and (1, 0, 0) is a vector that is orthogonal to all the simple roots. Thus these vectors are interchangeable. But for convenience avoiding fractional weights is advantageous.

However if you want to be an SL purist, Sage will support you. The weight space for SL(3) can be taken to be the hyperplane in ${\bf Q}^3$ consisting of vectors (a,b,c) with a+b+c=0. The fundamental weights for SL(3) are then (2/3,-1/3,-1/3) and (1/3,1/3,-2/3), though Sage will tell you they are (1,0,0) and (1,1,0). The work-around is to filter them through the method <code>coerce_to_sl</code> as follows:

```
sage: A2 = WeylCharacterRing("A2")
sage: [fw1,fw2] = [w.coerce_to_sl() for w in A2.fundamental_weights()]
```

```
sage: [standard, contragredient] = [A2(fw1), A2(fw2)]
sage: standard, contragredient
(A2(2/3, -1/3, -1/3), A2(1/3, 1/3, -2/3))
sage: standard*contragredient
A2(0,0,0) + A2(1,0,-1)
```

Sage is not confused by the fractional weights. Note that if you use coroot notation, you are working with SL automatically:

```
sage: A2 = WeylCharacterRing("A2", style="coroots")
sage: A2(1,0).weight_multiplicities()
{(-1/3, -1/3, 2/3): 1, (-1/3, 2/3, -1/3): 1, (2/3, -1/3, -1/3): 1}
```

There is no convenient way to create the determinant in the Weyl character ring if you adopt the coroot style.

Just as we coerced the fundamental weights into the SL weight lattice, you may need to coerce the Weyl vector ρ if you are working with SL. The default value for ρ in type A_r is $(r, r-1, \ldots, 0)$, but if you are an SL purist you want

$$\left(\frac{r}{2},\frac{r}{2}-1,\ldots,-\frac{r}{2}\right).$$

Therefore take the value of ρ that you get from the method of the ambient space and coerce it into SL:

```
sage: A2 = WeylCharacterRing("A2", style="coroots")
sage: rho = A2.space().rho().coerce_to_sl(); rho
(1, 0, -1)
sage: rho == (1/2)*sum(A2.space().positive_roots())
True
```

You do not need to do this for other Cartan types. If you are working with (say) F4 then a ρ is a ρ :

```
sage: F4 = WeylCharacterRing("F4")
sage: L = F4.space()
sage: rho = L.rho()
sage: rho == (1/2)*sum(L.positive_roots())
True
```

Integration

Suppose that we wish to compute the integral

$$\int_{U(n)} |tr(g)|^{2k} \, dg$$

for various n. Here U(n) is the unitary group, which is the maximal compact subroup of $GL(n, \mathbf{C})$, and dg is the Haar measure on U(n), normalized so that the volume of the group is 1.

The irreducible unitary representations of U(n) may be regarded as the basis elements of the WeylCharacterRing of type A_r , where r=n-1 so we might work in that ring. The trace tr(g) is then just the character of the standard representation. We may realize it in the WeylCharacterRing by taking the first fundamental weight and coercing it into the ring. For example, if k=5 and n=3 so r=2:

```
sage: A2 = WeylCharacterRing("A2")
sage: fw = A2.fundamental_weights(); fw
Finite family {1: (1, 0, 0), 2: (1, 1, 0)}
sage: tr = A2(fw[1]); tr
A2(1,0,0)
```

We may compute the norm square the character tr^5 by decomposing it into irreducibles, and taking the sum of the squares of their multiplicities. By Schur orthogonality, this gives the inner product of the $tr(g)^5$ with itself, that is, the integral of $|tr(g)|^{10}$:

```
sage: tr^5
5*A2(2,2,1) + 6*A2(3,1,1) + 5*A2(3,2,0) + 4*A2(4,1,0) + A2(5,0,0)
sage: sorted((tr^5).monomials(), key=lambda x: x.support())
[A2(2,2,1), A2(3,1,1), A2(3,2,0), A2(4,1,0), A2(5,0,0)]
sage: sorted((tr^5).coefficients())
[1, 4, 5, 5, 6]
sage: sum(x^2 for x in (tr^5).coefficients())
103
```

So far we have been working with n = 3. For general n:

```
sage: def f(n,k):
....: R = WeylCharacterRing(['A',n-1])
....: tr = R(R.fundamental_weights()[1])
....: return sum(x^2 for x in (tr^k).coefficients())
....:
sage: [f(n,5) for n in [2..7]] # long time (31s on sage.math, 2012)
[42, 103, 119, 120, 120, 120]
```

We see that the 10-th moment of tr(g) is just 5! when n is sufficiently large. What if we fix n and vary k?

```
sage: [f(2,k) for k in [1..10]]
[1, 2, 5, 14, 42, 132, 429, 1430, 4862, 16796]
sage: [catalan_number(k) for k in [1..10]]
[1, 2, 5, 14, 42, 132, 429, 1430, 4862, 16796]
```

Invariants and multiplicities

Sometimes we are only interested in the multiplicity of the trivial representation in some character. This may be found by the method invariant_degree. Continuing from the preceding example,

```
sage: A2 = WeylCharacterRing("A2", style="coroots")
sage: ad = A2(1,1)
sage: [ad.symmetric_power(k).invariant_degree() for k in [0..6]]
[1, 0, 1, 1, 1, 1, 2]
sage: [ad.exterior_power(k).invariant_degree() for k in [0..6]]
[1, 0, 0, 1, 0, 1, 0]
```

If we want the multiplicity of some other representation, we may obtain that using the method multiplicity:

```
sage: (ad^3).multiplicity(ad)
```

Maximal Subgroups and Branching Rules

Branching rules

If G is a Lie group and H is a subgroup, one often needs to know how representations of G restrict to H. Irreducibles usually do not restrict to irreducibles. In some cases the restriction is regular and predictable, in other cases it is chaotic. In some cases it follows a rule that can be described combinatorially, but the combinatorial description is subtle. The description of how irreducibles decompose into irreducibles is called a *branching rule*.

References for this topic:

- [FauserEtAl2006]
- [King1975]
- [HoweEtAl2005]
- [McKayPatera1981]

Sage can compute how a character of G restricts to H. It does so not by memorizing a combinatorial rule, but by computing the character and restricting the character to a maximal torus of H. What Sage has memorized (in a series of built-in encoded rules) are the various embeddings of maximal tori of maximal subgroups of G. The maximal subgroups of Lie groups were determined in [Dynkin1952]. This approach to computing branching rules has a limitation: the character must fit into memory and be computable by Sage's internal code in real time.

It is sufficient to consider the case where H is a maximal subgroup of G, since if this is known then one may branch down successively through a series of subgroups, each maximal in its predecessors. A problem is therefore to understand the maximal subgroups in a Lie group, and to give branching rules for each, and a goal of this tutorial is to explain the embeddings of maximal subgroups.

Sage has a class BranchingRule for branching rules. The function branching_rule returns elements of this class. For example, the natural embedding of Sp(4) into SL(4) corresponds to the branching rule that we may create as follows:

```
sage: b=branching_rule("A3","C2",rule="symmetric"); b
symmetric branching rule A3 => C2
```

The name "symmetric" of this branching rule will be explained further later, but it means that Sp(4) is the fixed subgroup of an involution of Sl(4). Here A3 and C2 are the Cartan types of the groups G = SL(4) and H = Sp(4).

Now we may see how representations of SL(4) decompose into irreducibles when they are restricted to Sp(4):

```
sage: A3=WeylCharacterRing("A3",style="coroots")
sage: chi=A3(1,0,1); chi.degree()
15
sage: C2=WeylCharacterRing("C2",style="coroots")
sage: chi.branch(C2,rule=b)
C2(0,1) + C2(2,0)
```

Alternatively, we may pass chi to b as an argument of its branch method, which gives the same result:

```
sage: b.branch(chi)
C2(0,1) + C2(2,0)
```

It is believed that the built-in branching rules of Sage are sufficient to handle all maximal subgroups and this is certainly the case when the rank if less than or equal to 8.

However, if you want to branch to a subgroup that is not maximal you may not find a built-in branching rule. We may compose branching rules to build up embeddings. For example, here are two different embeddings of Sp(4) with Cartan type C2 in Sp(8), with Cartan type C4. One embedding factors through $Sp(4) \times Sp(4)$, while the other factors through Sp(4). To check that the embeddings are not conjugate, we branch a (randomly chosen) representation. Observe that we do not have to build the intermediate Weyl character rings.

```
sage: C4=WeylCharacterRing("C4", style="coroots")
sage: b1=branching_rule("C4", "A3", "levi") *branching_rule("A3", "C2", "symmetric"); b1
composite branching rule C4 => (levi) A3 => (symmetric) C2
sage: b2=branching_rule("C4", "C2xC2", "orthogonal_sum") *branching_rule("C2xC2", "C2", "proj1"); b2
composite branching rule C4 => (orthogonal_sum) C2xC2 => (proj1) C2
sage: C2=WeylCharacterRing("C2", style="coroots")
sage: C4=WeylCharacterRing("C4", style="coroots")
```

```
sage: [C4(2,0,0,1).branch(C2, rule=br) for br in [b1,b2]] [4*C2(0,0) + 7*C2(0,1) + 15*C2(2,0) + 7*C2(0,2) + 11*C2(2,1) + C2(0,3) + 6*C2(4,0) + 3*C2(2,2), 10*C2(0,0) + 40*C2(1,0) + 50*C2(0,1) + 16*C2(2,0) + 20*C2(1,1) + 4*C2(3,0) + 5*C2(2,1)]
```

What's in a branching rule?

The essence of the branching rule is a function from the weight lattice of G to the weight lattice of the subgroup H, usually implemented as a function on the ambient vector spaces. Indeed, we may conjugate the embedding so that a Cartan subalgebra U of H is contained in a Cartan subalgebra T of G. Since the ambient vector space of the weight lattice of G is $\operatorname{Lie}(T)^*$, we get map $\operatorname{Lie}(T)^* \to \operatorname{Lie}(U)^*$, and this must be implemented as a function. For speed, the function usually just returns a list, which can be coerced into $\operatorname{Lie}(U)^*$.

```
sage: b = branching_rule("A3","C2","symmetric")
sage: for r in RootSystem("A3").ambient_space().simple_roots(): print r, b(r)
(1, -1, 0, 0) [1, -1]
(0, 1, -1, 0) [0, 2]
(0, 0, 1, -1) [1, -1]
```

We could conjugate this map by an element of the Weyl group of G, and the resulting map would give the same decomposition of representations of G into irreducibles of H. However it is a good idea to choose the map so that it takes dominant weights to dominant weights, and, insofar as possible, simple roots of G into simple roots of G. This includes sometimes the affine root G0 of G0, which we recall is the negative of the highest root.

The branching rule has a describe () method that shows how the roots (including the affine root) restrict. This is a useful way of understanding the embedding. You might want to try it with various branching rules of different kinds, "extended", "symmetric", "levi" etc.

```
sage: b.describe()
0
0----+
--0---0
    2
1
A3~
root restrictions A3 => C2:
\bigcirc = < = \bigcirc
1
    2.
C2
1 => 1
2 => 2
3 => 1
For more detailed information use verbose=True
```

The extended Dynkin diagram of G and the ordinary Dynkin diagram of H are shown for reference, and 3 > 1 means that the third simple root α_3 of G restricts to the first simple root of H. In this example, the affine root does not restrict to a simple roots, so it is omitted from the list of restrictions. If you add the parameter verbose=true you will be shown the restriction of all simple roots and the affine root, and also the restrictions of the fundamental weights (in coroot notation).

Maximal subgroups

Sage has a database of maximal subgroups for every simple Cartan type of rank ≤ 8 . You may access this with the maximal_subgroups method of the Weyl character ring:

```
sage: E7=WeylCharacterRing("E7",style="coroots")
sage: E7.maximal_subgroups()
A7:branching_rule("E7","A7","extended")
E6:branching_rule("E7","E6","levi")
A2:branching_rule("E7","A2","miscellaneous")
A1:branching_rule("E7","A1","iii")
A1:branching_rule("E7","A1","iv")
A1xF4:branching_rule("E7","A1xF4","miscellaneous")
G2xC3:branching_rule("E7","G2xC3","miscellaneous")
A1xG2:branching_rule("E7","A1xG2","miscellaneous")
A1xA1:branching_rule("E7","A1xA1","miscellaneous")
A1xD6:branching_rule("E7","A1xD6","extended")
A5xA2:branching_rule("E7","A5xA2","extended")
```

It should be understood that there are other ways of embedding $A_2 = SL(3)$ into the Lie group E_7 , but only one way as a maximal subgroup. On the other hand, there are but only one way to embed it as a maximal subgroup. The embedding will be explained below. You may obtain the branching rule as follows, and use it to determine the decomposition of irreducible representations of E_7 as follows:

```
sage: b = E7.maximal_subgroup("A2"); b
miscellaneous branching rule E7 => A2
sage: [E7,A2]=[WeylCharacterRing(x,style="coroots") for x in ["E7","A2"]]
sage: E7(1,0,0,0,0,0,0).branch(A2,rule=b)
A2(1,1) + A2(4,4)
```

This gives the same branching rule as just pasting line beginning to the right of the colon onto the command line:

```
sage:branching_rule("E7","A2","miscellaneous")
miscellaneous branching rule E7 => A2
```

There are two distict embeddings of $A_1 = SL(2)$ into E_7 as maximal subgroups, so the maximal_subgroup method will return a list of rules:

```
sage: WeylCharacterRing("E7").maximal_subgroup("A1")
[iii branching rule E7 => A1, iv branching rule E7 => A1]
```

The list of maximal subgroups returned by the maximal_subgroups method for irreducible Cartan types of rank up to 8 is believed to be complete up to outer automorphisms. You may want a list that is complete up to inner automorphisms. For example, E_6 has a nontrivial Dynkin diagram automorphism so it has an outer automorphism that is not inner:

```
sage: [E6,A2xG2]=[WeylCharacterRing(x,style="coroots") for x in ["E6","A2xG2"]]
sage: b=E6.maximal_subgroup("A2xG2"); b
miscellaneous branching rule E6 => A2xG2
sage: E6(1,0,0,0,0,0).branch(A2xG2,rule=b)
A2xG2(0,1,1,0) + A2xG2(2,0,0,0)
sage: E6(0,0,0,0,0,1).branch(A2xG2,rule=b)
A2xG2(1,0,1,0) + A2xG2(0,2,0,0)
```

Since as we see the two 27 dimensional irreducibles (which are interchanged by the outer automorphism) have different branching, the $A_2 \times G_2$ subgroup is changed to a different one by the outer automorphism. To obtain the second branching rule, we compose the given one with this automorphism:

```
sage: b1=branching_rule("E6","E6","automorphic")*b; b1
composite branching rule E6 => (automorphic) E6 => (miscellaneous) A2xG2
```

Levi subgroups

A Levi subgroup may or may not be maximal. They are easily classified. If one starts with a Dynkin diagram for G and removes a single node, one obtains a smaller Dynkin diagram, which is the Dynkin diagram of a smaller subgroup H.

For example, here is the A3 Dynkin diagram:

```
sage: A3 = WeylCharacterRing("A3")
sage: A3.dynkin_diagram()
0---0--0
1          2          3
A3
```

We see that we may remove the node 3 and obtain A_2 , or the node 2 and obtain $A_1 \times A_1$. These correspond to the Levi subgroups GL(3) and $GL(2) \times GL(2)$ of GL(4).

Let us construct the irreducible representations of GL(4) and branch them down to these down to GL(3) and $GL(2) \times GL(2)$:

```
sage: reps = [A3(v) for v in A3.fundamental_weights()]; reps
[A3(1,0,0,0), A3(1,1,0,0), A3(1,1,1,0)]
sage: A2 = WeylCharacterRing("A2")
sage: A1xA1 = WeylCharacterRing("A1xA1")
sage: [pi.branch(A2, rule="levi") for pi in reps]
[A2(0,0,0) + A2(1,0,0), A2(1,0,0) + A2(1,1,0), A2(1,1,0) + A2(1,1,1)]
sage: [pi.branch(A1xA1, rule="levi") for pi in reps]
[A1xA1(1,0,0,0) + A1xA1(0,0,1,0),
A1xA1(1,1,0,0) + A1xA1(1,0,1,0) + A1xA1(0,0,1,1),
A1xA1(1,1,1,0) + A1xA1(1,0,1,1)]
```

Let us redo this calculation in coroot notation. As we have explained, coroot notation does not distinguish between representations of GL(4) that have the same restriction to SL(4), so in effect we are now working with the groups SL(4) and its Levi subgroups SL(3) and $SL(2) \times SL(2)$, which is the derived group of its Levi subgroup:

```
sage: A3 = WeylCharacterRing("A3", style="coroots")
sage: reps = [A3(v) for v in A3.fundamental_weights()]; reps
[A3(1,0,0), A3(0,1,0), A3(0,0,1)]
sage: A2 = WeylCharacterRing("A2", style="coroots")
sage: A1xA1 = WeylCharacterRing("A1xA1", style="coroots")
sage: [pi.branch(A2, rule="levi") for pi in reps]
[A2(0,0) + A2(1,0), A2(0,1) + A2(1,0), A2(0,0) + A2(0,1)]
sage: [pi.branch(A1xA1, rule="levi") for pi in reps]
[A1xA1(1,0) + A1xA1(0,1), 2*A1xA1(0,0) + A1xA1(1,1), A1xA1(1,0) + A1xA1(0,1)]
```

Now we may observe a distinction difference in branching from

$$GL(4) \to GL(2) \times GL(2)$$

versus

$$SL(4) \rightarrow SL(2) \times SL(2)$$
.

Consider the representation A3 (0, 1, 0), which is the six dimensional exterior square. In the coroot notation, the restriction contained two copies of the trivial representation, $2 * A1 \times A1 (0, 0)$. The other way, we had instead three

distinct representations in the restriction, namely A1xA1 (1,1,0,0) and A1xA1 (0,0,1,1), that is, $\det \otimes 1$ and $1 \otimes \det$.

The Levi subgroup A1xA1 is actually not maximal. Indeed, we may factor the embedding:

$$SL(2) \times SL(2) \rightarrow Sp(4) \rightarrow SL(4)$$
.

Therfore there are branching rules A3 -> C2 and C2 -> A2, and we could accomplish the branching in two steps, thus:

As you can see, we've redone the branching rather circuitously this way, making use of the branching rules A3 -> C2 and B2 -> D2, and two accidental isomorphisms C2 = B2 and D2 = A1xA1. It is much easier to go in one step using rule="levi", but reassuring that we get the same answer!

Subgroups classified by the extended Dynkin diagram

It is also true that if we remove one node from the extended Dynkin diagram that we obtain the Dynkin diagram of a subgroup. For example:

```
sage: G2 = WeylCharacterRing("G2", style="coroots")
sage: G2.extended_dynkin_diagram()
    3
0=<=0---0
1    2    0
G2~</pre>
```

Observe that by removing the 1 node that we obtain an A_2 Dynkin diagram. Therefore the exceptional group G_2 contains a copy of SL(3). We branch the two representations of G_2 corresponding to the fundamental weights to this copy of A_2 :

```
sage: G2 = WeylCharacterRing("G2", style="coroots")
sage: A2 = WeylCharacterRing("A2", style="coroots")
sage: [G2(f).degree() for f in G2.fundamental_weights()]
[7, 14]
sage: [G2(f).branch(A2, rule="extended") for f in G2.fundamental_weights()]
[A2(0,0) + A2(0,1) + A2(1,0), A2(0,1) + A2(1,0) + A2(1,1)]
```

The two representations of G_2 , of degrees 7 and 14 respectively, are the action on the octonions of trace zero and the adjoint representation.

For embeddings of this type, the rank of the subgroup H is the same as the rank of G. This is in contrast with embeddings of Levi type, where H has rank one less than G.

Levi subgroups of G_2

The exceptional group G_2 has two Levi subgroups of type A_1 . Neither is maximal, as we can see from the extended Dynkin diagram: the subgroups $A_1 \times A_1$ and A_2 are maximal and each contains a Levi subgroup. (Actually $A_1 \times A_1$

contains a conjugate of both.) Only the Levi subgroup containing the short root is implemented as an instance of rule="levi". To obtain the other, use the rule:

```
sage: branching_rule("G2", "A2", "extended") *branching_rule("A2", "A1", "levi")
composite branching rule G2 => (extended) A2 => (levi) A1
```

which branches to the A_1 Levi subgroup containing a long root.

Orthogonal and symplectic subgroups of orthogonal and symplectic groups

If G = SO(n) then G has a subgroup SO(n-1). Depending on whether n is even or odd, we thus have branching rules ['D', r] to ['B', r-1] or ['B', r] to ['D', r]. These are handled as follows:

```
sage: branching_rule("B4","D4",rule="extended")
extended branching rule B4 => D4
sage: branching_rule("D4","B3",rule="symmetric")
symmetric branching rule D4 => B3
```

If G = SO(r + s) then G has a subgroup $SO(r) \times SO(s)$. This lifts to an embedding of the universal covering groups

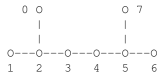
$$spin(r) \times spin(s) \rightarrow spin(r+s)$$
.

Sometimes this embedding is of extended type, and sometimes it is not. It is of extended type unless r and s are both odd. If it is of extended type then you may use rule="extended". In any case you may use rule="orthogonal_sum". The name refer to the origin of the embedding $SO(r) \times SO(s) \rightarrow SO(r+s)$ from the decomposition of the underlying quadratic space as a direct sum of two orthogonal subspaces.

There are four cases depending on the parity of r and s. For example, if r = 2k and s = 2l we have an embedding:

```
['D',k] x ['D',1] --> ['D',k+1]
```

This is of extended type. Thus consider the embedding $D4 \times D3 -> D7$. Here is the extended Dynkin diagram:



Removing the 4 vertex results in a disconnected Dynkin diagram:

This is D4xD3. Therefore use the "extended" branching rule:

```
sage: D7 = WeylCharacterRing("D7", style="coroots")
sage: D4xD3 = WeylCharacterRing("D4xD3", style="coroots")
sage: spin = D7(D7.fundamental_weights()[7]); spin
D7(0,0,0,0,0,0,1)
sage: spin.branch(D4xD3, rule="extended")
D4xD3(0,0,1,0,0,1,0) + D4xD3(0,0,0,1,0,0,1)
```

But we could equally well use the "orthogonal_sum" rule:

```
sage: spin.branch(D4xD3, rule="orthogonal_sum")
D4xD3(0,0,1,0,0,1,0) + D4xD3(0,0,0,1,0,0,1)
```

Similarly we have embeddings:

```
['D',k] x ['B',1] --> ['B',k+1]
```

These are also of extended type. For example consider the embedding of D3xB2 -> B5. Here is the B5 extended Dynkin diagram:

```
0 0
|
|----0--0--0=>=0
1 2 3 4 5
```

Removing the 3 node gives:

and this is the Dynkin diagram or D3xB2. For such branchings we again use either rule="extended" or rule="orthogonal_sum".

Finally, there is an embedding

```
['B',k] x ['B',1] --> ['D',k+1+1]
```

This is not of extended type, so you may not use rule="extended". You must use rule="orthogonal_sum":

```
sage: D5 = WeylCharacterRing("D5",style="coroots")
sage: B2xB2 = WeylCharacterRing("B2xB2",style="coroots")
sage: [D5(v).branch(B2xB2,rule="orthogonal_sum") for v in D5.fundamental_weights()]
[B2xB2(1,0,0,0) + B2xB2(0,0,1,0),
B2xB2(0,2,0,0) + B2xB2(1,0,1,0) + B2xB2(0,0,0,2),
B2xB2(0,2,0,0) + B2xB2(0,2,1,0) + B2xB2(1,0,0,2) + B2xB2(0,0,0,2),
B2xB2(0,1,0,1), B2xB2(0,1,0,1)]
```

Non-maximal Levi subgroups and Projection from Reducible Types

Not all Levi subgroups are maximal. Recall that the Dynkin-diagram of a Levi subgroup H of G is obtained by removing a node from the Dynkin diagram of G. Removing the same node from the extended Dynkin diagram of G results in the Dynkin diagram of a subgroup of G that is strictly larger than H. However this subgroup may or may not be proper, so the Levi subgroup may or may not be maximal.

If the Levi subgroup is not maximal, the branching rule may or may not be implemented in Sage. However if it is not implemented, it may be constructed as a composition of two branching rules.

For example, prior to Sage-6.1 branching_rule ("E6", "A5", "levi") returned a not-implemented error and the advice to branch to A5xA1. And we can see from the extended Dynkin diagram of E_6 that indeed A_5 is not a maximal subgroup, since removing node 2 from the extended Dynkin diagram (see below) gives A5xA1. To construct the branching rule to A_5 we may proceed as follows:

```
sage: b = branching_rule("E6", "A5xA1", "extended") *branching_rule("A5xA1", "A5", "proj1"); b
composite branching rule E6 => (extended) A5xA1 => (proj1) A5
sage: E6=WeylCharacterRing("E6", style="coroots")
sage: A5=WeylCharacterRing("A5", style="coroots")
sage: E6(0,1,0,0,0,0).branch(A5,rule=b)
3*A5(0,0,0,0,0) + 2*A5(0,0,1,0,0) + A5(1,0,0,0,1)
sage: b.describe()
       0 0
       0 2
        0---0---0
   3 4 5 6
1
E6~
root restrictions E6 => A5:
0---0
1
   2 3 4 5
A 5
0 \Rightarrow (zero)
1 => 1
3 => 2
4 => 3
5 => 4
6 => 5
```

Note that it is not necessary to construct the Weyl character ring for the intermediate group A5xA1.

This last example illustrates another common problem: how to extract one component from a reducible root system. We used the rule "proj1" to extract the first component. We could similarly use "proj2" to get the second, or more generally any combination of components:

```
sage: branching_rule("A2xB2xG2","A2xG2","proj13")
proj13 branching rule A2xB2xG2 => A2xG2
```

For more detailed information use verbose=True

Symmetric subgroups

If G admits an outer automorphism (usually of order two) then we may try to find the branching rule to the fixed subgroup H. It can be arranged that this automorphism maps the maximal torus T to itself and that a maximal torus U of H is contained in T.

Suppose that the Dynkin diagram of G admits an automorphism. Then G itself admits an outer automorphism. The Dynkin diagram of the group H of invariants may be obtained by "folding" the Dynkin diagram of G along the automorphism. The exception is the branching rule $GL(2r) \to SO(2r)$.

Here are the branching rules that can be obtained using rule="symmetric".

G	Н	Cartan Types
GL(2r)	Sp(2r)	['A',2r-1] => ['C',r]
GL(2r+1)	SO(2r+1)	['A',2r] => ['B',r]
GL(2r)	SO(2r)	['A',2r-1] => ['D',r]
SO(2r)	SO(2r-1)	['D',r] => ['B',r-1]
E_6	F_4	['E',6] => ['F',4]

Tensor products

If G_1 and G_2 are Lie groups, and we have representations $\pi_1: G_1 \to GL(n)$ and $\pi_2: G_2 \to GL(m)$ then the tensor product is a representation of $G_1 \times G_2$. It has its image in GL(nm) but sometimes this is conjugate to a subgroup of SO(nm) or Sp(nm). In particular we have the following cases.

Group	Subgroup	Cartan Types
GL(rs)	$GL(r) \times GL(s)$	$['A', rs-1] \Rightarrow ['A', r-1] \times ['A', s-1]$
SO(4rs + 2r + 2s + 1)	$SO(2r+1) \times SO(2s+1)$	['B',2rs+r+s] => ['B',r] x ['B',s]
SO(4rs + 2s)	$SO(2r+1) \times SO(2s)$	['D',2rs+s] => ['B',r] x ['D',s]
SO(4rs)	$SO(2r) \times SO(2s)$	['D',2rs] => ['D',r] x ['D',s]
SO(4rs)	$Sp(2r) \times Sp(2s)$	['D',2rs] => ['C',r] x ['C',s]
Sp(4rs + 2s)	$SO(2r+1) \times Sp(2s)$	['C',2rs+s] => ['B',r] x ['C',s]
Sp(4rs)	$Sp(2r) \times SO(2s)$	['C',2rs] => ['C',r] x ['D',s]

These branching rules are obtained using rule="tensor".

Symmetric powers

The k-th symmetric and exterior power homomorphisms map $GL(n) \to GL\left(\binom{n+k-1}{k}\right)$ and $GL\left(\binom{n}{k}\right)$. The corresponding branching rules are not implemented but a special case is. The k-th symmetric power homomorphism $SL(2) \to GL(k+1)$ has its image inside of SO(2r+1) if k=2r and inside of Sp(2r) if k=2r-1. Hence there are branching rules:

and these may be obtained using rule="symmetric_power".

Plethysms

The above branching rules are sufficient for most cases, but a few fall between the cracks. Mostly these involve maximal subgroups of fairly small rank.

The rule rule="plethysm" is a powerful rule that includes any branching rule from types A, B, C or D as a special case. Thus it could be used in place of the above rules and would give the same results. However, it is most useful when branching from G to a maximal subgroup H such that rank(H) < rank(G) - 1.

We consider a homomorphism $H \to G$ where G is one of SL(r+1), SO(2r+1), Sp(2r) or SO(2r). The function branching_rule_from_plethysm produces the corresponding branching rule. The main ingredient is the character χ of the representation of H that is the homomorphism to GL(r+1), GL(2r+1) or GL(2r).

Let us consider the symmetric fifth power representation of SL(2). This is implemented above by rule="symmetric_power", but suppose we want to use rule="plethysm". First we construct the homomorphism by invoking its character, to be called chi:

```
sage: A1 = WeylCharacterRing("A1", style="coroots")
sage: chi = A1([5])
sage: chi.degree()
6
sage: chi.frobenius_schur_indicator()
-1
```

This confirms that the character has degree 6 and is symplectic, so it corresponds to a homomorphism $SL(2) \to Sp(6)$, and there is a corresponding branching rule C3 -> A1:

```
sage: A1 = WeylCharacterRing("A1", style="coroots")
sage: C3 = WeylCharacterRing("C3", style="coroots")
sage: chi = A1([5])
sage: sym5rule = branching_rule_from_plethysm(chi, "C3")
sage: [C3(hwv).branch(A1, rule=sym5rule) for hwv in C3.fundamental_weights()]
[A1(5), A1(4) + A1(8), A1(3) + A1(9)]
```

This is identical to the results we would obtain using rule="symmetric_power":

```
sage: A1 = WeylCharacterRing("A1", style="coroots")
sage: C3 = WeylCharacterRing("C3", style="coroots")
sage: [C3(v).branch(A1, rule="symmetric_power") for v in C3.fundamental_weights()]
[A1(5), A1(4) + A1(8), A1(3) + A1(9)]
```

But the next example of plethysm gives a branching rule not available by other methods:

```
sage: G2 = WeylCharacterRing("G2", style="coroots")
sage: D7 = WeylCharacterRing("D7", style="coroots")
sage: ad = G2.adjoint_representation(); ad.degree()
14
sage: ad.frobenius_schur_indicator()
1
sage: for r in D7.fundamental_weights(): # long time (1.29s)
....: print D7(r).branch(G2, rule=branching_rule_from_plethysm(ad, "D7"))
....:
G2(0,1)
G2(0,1) + G2(3,0)
G2(0,0) + G2(2,0) + G2(3,0) + G2(0,2) + G2(4,0)
G2(0,1) + G2(2,0) + G2(1,1) + G2(0,2) + G2(2,1) + G2(4,0) + G2(3,1)
G2(1,0) + G2(0,1) + G2(1,1) + 2*G2(3,0) + 2*G2(2,1) + G2(1,2) + G2(3,1) + G2(5,0) + G2(0,3)
G2(1,1)
G2(1,1)
```

In this example, ad is the 14-dimensional adjoint representation of the exceptional group G_2 . Since the Frobenius-Schur indicator is 1, the representation is orthogonal, and factors through SO(14), that is, D7.

We do not actually have to create the character (or for that matter its ambient WeylCharacterRing) in order to create the branching rule:

```
sage: branching_rule("D4", "A2.adjoint_representation()", "plethysm")
plethysm (along A2(1,1)) branching rule D4 => A2
```

The adjoint representation of any semisimple Lie group is orthogonal, so we do not need to compute the Frobenius-Schur indicator.

Miscellaneous other subgroups

Use rule="miscellaneous" for the following rules. Every maximal subgroup H of an exceptional group G are either among these, or the five A_1 subgroups described in the next section, or (if G and H have the same rank) is available using rule="extended".

$$\begin{split} &B_3 \to G_2, \\ &E_6 \to A_2, \\ &E_6 \to G_2, \\ &F_4 \to G_2 \times A_1, \\ &E_6 \to G_2 \times A_2, \\ &E_7 \to G_2 \times C_3, \\ &E_7 \to F_4 \times A_1, \\ &E_7 \to A_1 \times A_1, \\ &E_7 \to G_2 \times A_1, \\ &E_7 \to A_2 \\ &E_8 \to G_2 \times F_4. \\ &E_8 \to A_2 \times A_1. \\ &E_8 \to B_2 \end{split}$$

The first rule corresponds to the embedding of G_2 in SO(7) in its action on the trace zero octonions. The two branching rules from E_6 to G_2 or A_2 are described in [Testerman1989]. We caution the reader that Theorem G.2 of that paper, proved there in positive characteristic is false over the complex numbers. On the other hand, the assumption of characteristic p is not important for Theorems G.1 and A.1, which describe the torus embeddings, hence contain enough information to compute the branching rule. There are other ways of embedding G_2 or G_2 into G_3 or G_4 into G_4 into G_4 or G_4 into G_4 into G_4 or G_4 into G_4 into

The remaining rules come about as follows. Let G be F_4 , E_6 , E_7 or E_8 , and let H be G_2 , or else (if $G = E_7$) F_4 . We embed H into G in the most obvious way; that is, in the chain of subgroups

$$G_2 \subset F_4 \subset E_6 \subset E_7 \subset E_8$$

Then the centralizer of H is A_1 , A_2 , C_3 , F_4 (if $H = G_2$) or A_1 (if $G = E_7$ and $H = F_4$). This gives us five of the cases. Regarding the branching rule $E_6 \to G_2 \times A_2$, Rubenthaler [Rubenthaler2008] describes the embedding and applies it in an interesting way.

The embedding of $A_1 \times A_1$ into E_7 is as follows. Deleting the 5 node of the E_7 Dynkin diagram gives the Dynkin diagram of $A_4 \times A_2$, so this is a Levi subgroup. We embed SL(2) into this Levi subgroup via the representation $[4] \otimes [2]$. This embeds the first copy of A_1 . The other A_1 is the connected centralizer. See [Seitz1991], particularly the proof of (3.12).

The embedding if $G_2 \times A_1$ into E_7 is as follows. Deleting the 2 node of the E_7 Dynkin diagram gives the A_6 Dynkin diagram, which is the Levi subgroup SL(7). We embed G_2 into SL(7) via the irreducible seven-dimensional representation of G_2 . The A_1 is the centralizer.

The embedding if $A_2 \times A_1$ into E_8 is as follows. Deleting the 2 node of the E_8 Dynkin diagram gives the A_7 Dynkin diagram, which is the Levi subgroup SL(8). We embed A_2 into SL(8) via the irreducible eight-dimensional adjoint representation of SL(2). The A_1 is the centralizer.

The embedding A_2 into E_7 is proved in [Seitz1991] (5.8). In particular, he computes the embedding of the SL(3) torus in the E_7 torus, which is what is needed to implement the branching rule. The embedding of B_2 into E_8 is also constructed in [Seitz1991] (6.7). The embedding of the B_2 Cartan subalgebra, needed to implement the branching rule, is easily deduced from (10) on page 111.

Maximal A1 subgroups of Exceptional Groups

There are seven embeddings of SL(2) into an exceptional group as a maximal subgroup: one each for G_2 and F_4 , two nonconjugate embeddings for E_7 and three for E_8 These are constructed in [Testerman1992]. Create the corresponding branching rules as follows. The names of the rules are roman numerals referring to the seven cases of Testerman's Theorem 1:

```
sage: branching_rule("G2", "A1", "i")
i branching rule G2 => A1
sage: branching_rule("F4", "A1", "ii")
ii branching rule F4 => A1
sage: branching_rule("E7", "A1", "iii")
iii branching rule E7 => A1
sage: branching_rule("E7", "A1", "iv")
iv branching rule E7 => A1
sage: branching_rule("E8", "A1", "v")
v branching rule E8 => A1
sage: branching_rule("E8", "A1", "vi")
vi branching rule E8 => A1
sage: branching_rule("E8", "A1", "vi")
vi branching rule E8 => A1
```

The embeddings are characterized by the root restrictions in their branching rules: usually a simple root of the ambient group G restricts to the unique simple root of A_1 , except for root α_4 for rules iv, vi and vii, and the root α_6 for root vii; this is essentially the way Testerman characterizes the embeddings, and this information may be obtained from Sage by employing the describe () method of the branching rule. Thus:

```
sage: branching_rule("E8", "A1", "vii").describe()
        0 2
  --0---0---0---0
      4 5 6 7 8
1
    3
E.8~
root restrictions E8 => A1:
\bigcirc
1
A1
1 => 1
2 => 1
3 => 1
4 \Rightarrow (zero)
5 => 1
6 \Rightarrow (zero)
7 => 1
8 => 1
```

For more detailed information use verbose=True

Writing your own branching rules

Sage has many built-in branching rules. Indeed, at least up to rank eight (including all the exceptional groups) branching rules to all maximal subgroups are implemented as built in rules, except for a few obtainable using branching_rule_from_plethysm. This means that all the rules in [McKayPatera1981] are available in Sage.

Still in this section we are including instructions for coding a rule by hand. As we have already explained, the branching rule is a function from the weight lattice of G to the weight lattice of H, and if you supply this you can write your own branching rules.

As an example, let us consider how to implement the branching rule A3 \rightarrow C2. Here H = C2 = Sp (4) embedded as a subgroup in A3 = GL (4). The Cartan subalgebra Lie(U) consists of diagonal matrices with eigenvalues u1, u2, -u2, -u1. Then C2.space () is the two dimensional vector spaces consisting of the linear functionals u1 and u2 on U. On the other hand $Lie(T) = \mathbf{R}^4$. A convenient way to see the restriction is to think of it as the adjoint of the map [u1, u2] \rightarrow [u1, u2, -u2, -u1], that is, [x0, x1, x2, x3] \rightarrow [x0-x3, x1-x2]. Hence we may encode the rule:

```
def brule(x):
    return [x[0]-x[3], x[1]-x[2]]

or simply:
brule = lambda x: [x[0]-x[3], x[1]-x[2]]
```

Let us check that this agrees with the built-in rule:

```
sage: A3 = WeylCharacterRing(['A', 3])
sage: C2 = WeylCharacterRing(['C', 2])
sage: brule = lambda x: [x[0]-x[3], x[1]-x[2]]
sage: A3(1,1,0,0).branch(C2, rule=brule)
C2(0,0) + C2(1,1)
sage: A3(1,1,0,0).branch(C2, rule="symmetric")
C2(0,0) + C2(1,1)
```

Although this works, it is better to make the rule into an element of the BranchingRule class, as follows.

```
sage: brule = BranchingRule("A3", "C2", lambda x : [x[0]-x[3], x[1]-x[2]], "custom") sage: A3(1,1,0,0).branch(C2, rule=brule) C2(0,0) + C2(1,1)
```

Automorphisms and triality

The case where G = H can be treated as a special case of a branching rule. In most cases if G has a nontrivial outer automorphism, it has order two, corresponding to the symmetry of the Dynkin diagram. Such an involution exists in the cases A_r , D_r , E_6 .

So the automorphism acts on the representations of G, and its effect may be computed using the branching rule code:

```
sage: A4 = WeylCharacterRing("A4", style="coroots")
sage: A4(1,0,1,0).degree()
45
sage: A4(0,1,0,1).degree()
45
sage: A4(1,0,1,0).branch(A4,rule="automorphic")
A4(0,1,0,1)
```

In the special case where G=D4, the Dynkin diagram has extra symmetries, and these correspond to outer automorphisms of the group. These are implemented as the "triality" branching rule:

```
sage: branching_rule("D4", "D4", "triality").describe()
   0 4
   0---0
  12 3
   0 0
D4~
root restrictions D4 => D4:
   0 4
   0---0
1
   2
D4
1 => 3
2 = > 2
3 => 4
4 => 1
```

Triality his is not an automorphisms of SO(8), but of its double cover spin(8). Note that spin(8) has three representations of degree 8, namely the standard representation of SO(8) and the two eight-dimensional spin representations. These are permuted by triality:

```
sage: D4=WeylCharacterRing("D4",style="coroots")
sage: D4(0,0,0,1).branch(D4,rule="triality")
D4(1,0,0,0)
sage: D4(0,0,0,1).branch(D4,rule="triality").branch(D4,rule="triality")
D4(0,0,1,0)
sage: D4(0,0,0,1).branch(D4,rule="triality").branch(D4,rule="triality").branch(D4,rule="triality")
```

By contrast, rule="automorphic" simply interchanges the two spin representations, as it always does in type D:

```
sage: D4(0,0,0,1).branch(D4,rule="automorphic")
D4(0,0,1,0)
sage: D4(0,0,1,0).branch(D4,rule="automorphic")
D4(0,0,0,1)
```

For more detailed information use verbose=True

Weight Rings

You may wish to work directly with the weights of a representation.

Weyl character ring elements are represented internally by a dictionary of their weights with multiplicities. However these are subject to a constraint: the coefficients must be invariant under the action of the Weyl group.

The WeightRing is also a ring whose elements are represented internally by a dictionary of their weights with multiplicities, but it is not subject to this constraint of Weyl group invariance. The weights are allowed to be fractional,

that is, elements of the ambient space. In other words, the weight ring is the group algebra over the ambient space of the weight lattice.

To create a WeightRing first construct the WeylCharacterRing, then create the WeightRing as follows:

```
sage: A2 = WeylCharacterRing(['A',2])
sage: a2 = WeightRing(A2)
```

You may coerce elements of the WeylCharacterRing into the weight ring. For example, if you want to see the weights of the adjoint representation of GL(3), you may use the method mlist, but another way is to coerce it into the weight ring:

```
sage: from pprint import pprint
sage: A2 = WeylCharacterRing(['A',2])
sage: ad = A2(1,0,-1)
sage: pprint(ad.weight_multiplicities())
{(0, 0, 0): 2, (-1, 1, 0): 1, (-1, 0, 1): 1, (1, -1, 0): 1,
   (1, 0, -1): 1, (0, -1, 1): 1, (0, 1, -1): 1}
```

This command produces a dictionary of the weights that appear in the representation, together with their multiplicities. But another way of getting the same information, with an aim of working with it, is to coerce it into the weight ring:

```
sage: a2 = WeightRing(A2)
sage: a2(ad)
2*a2(0,0,0) + a2(-1,1,0) + a2(-1,0,1) + a2(1,-1,0) + a2(1,0,-1) + a2(0,-1,1) + a2(0,1,-1)
```

For example, the Weyl denominator formula is usually written this way:

$$\prod_{\alpha \in \Phi^+} \left(e^{\alpha/2} - e^{-\alpha/2} \right) = \sum_{w \in W} (-1)^{l(w)} e^{w(\rho)}.$$

The notation is as follows. Here if λ is a weight, or more generally, an element of the ambient space, then e^{λ} means the image of λ in the group algebra of the ambient space of the weight lattice λ . Since this group algebra is just the weight ring, we can interpret e^{λ} as its image in the weight ring.

Let us confirm the Weyl denominator formula for A2:

```
sage: A2 = WeylCharacterRing("A2")
sage: a2 = WeightRing(A2)
sage: L = A2.space()
sage: W = L.weyl_group()
sage: rho = L.rho().coerce_to_sl()
sage: lhs = prod(a2(alpha/2)-a2(-alpha/2) for alpha in L.positive_roots()); lhs
a2(-1,1,0) - a2(-1,0,1) - a2(1,-1,0) + a2(1,0,-1) + a2(0,-1,1) - a2(0,1,-1)
sage: rhs = sum((-1)^(w.length())*a2(w.action(rho)) for w in W); rhs
a2(-1,1,0) - a2(-1,0,1) - a2(1,-1,0) + a2(1,0,-1) + a2(0,-1,1) - a2(0,1,-1)
sage: lhs == rhs
True
```

Note that we have to be careful to use the right value of ρ . The reason for this is explained in *SL versus GL*.

We have seen that elements of the WeylCharacterRing can be coerced into the WeightRing. Elements of the WeightRing can be coerced into the WeylCharacterRing provided they are invariant under the Weyl group.

Weyl Groups, Coxeter Groups and the Bruhat Order

Classical and affine Weyl groups

You can create Weyl groups and affine Weyl groups for any root system. A variety of methods are available for these. Some of these are methods are available for general Coxeter groups.

By default, elements of the Weyl group are represented as matrices:

```
sage: WeylGroup("A3").simple_reflection(1)
[0 1 0 0]
[1 0 0 0]
[0 0 1 0]
[0 0 0 1]
```

You may prefer a notation in which elements are written out as products of simple reflections. In order to implement this you need to specify a prefix, typically "s":

```
sage: W = WeylGroup("A3",prefix="s")
sage: [s1,s2,s3] = W.simple_reflections()
sage: (s1*s2*s1).length()
3
sage: W.long_element()
s1*s2*s3*s1*s2*s1
sage: s1*s2*s3*s1*s2*s1 == s3*s2*s1*s3*s2*s3
```

The Weyl group acts on the ambient space of the root lattice, which is accessed by the method domain. To illustrate this, recall that if w_0 is the long element then $\alpha \mapsto -w_0(\alpha)$ is a permutation of the simple roots. We may compute this as follows:

```
sage: W = WeylGroup("E6",prefix="s")
sage: w0 = W.long_element(); w0
s1*s3*s4*s5*s6*s2*s4*s5*s3*s4*s1*s3*s2*s4*s5*s6*s2*s4*s5*s3*s4*s1*s3*s2*s4*s5*s3*s4*s1*s3*s2*s4*s1*s
sage: sr = W.domain().simple_roots().list(); sr
[(1/2, -1/2, -1/2, -1/2, -1/2, -1/2, -1/2, 1/2), (1, 1, 0, 0, 0, 0, 0, 0),
(-1, 1, 0, 0, 0, 0, 0, 0), (0, -1, 1, 0, 0, 0, 0, 0), (0, 0, -1, 1, 0, 0, 0, 0),
(0, 0, 0, -1, 1, 0, 0, 0)]
sage: [-w0.action(a) for a in sr]
[(0, 0, 0, -1, 1, 0, 0, 0), (1, 1, 0, 0, 0, 0, 0, 0), (0, 0, -1, 1, 0, 0, 0, 0),
(0, -1, 1, 0, 0, 0, 0, 0, 0), (-1, 1, 0, 0, 0, 0, 0),
(1/2, -1/2, -1/2, -1/2, -1/2, -1/2, -1/2, 1/2)]
```

We may ask when this permutation is trivial. If it is nontrivial it induces an automorphism of the Dynkin diagram, so it must be nontrivial when the Dynkin diagram has no automorphism. But if there is a nontrivial automorphism, the permutation might or might not be trivial:

```
sage: def roots_not_permuted(ct):
....:     W = WeylGroup(ct)
....:     w0 = W.long_element()
....:     sr = W.domain().simple_roots()
....:     return all(a == -w0.action(a) for a in sr)
....:
sage: for ct in [CartanType(['D',r]) for r in [2..8]]:
....:     print ct,roots_not_permuted(ct)
....:
['D', 2] True
['D', 3] False
```

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```
['D', 4] True
['D', 5] False
['D', 6] True
['D', 7] False
['D', 8] True
```

If α is a root let r_{α} denote the reflection in the hyperplane that is orthogonal to α . We reserve the notation s_{α} for the simple reflections, that is, the case where α is a simple root. The reflections are just the conjugates of the simple reflections.

The reflections are the keys in a finite family, which is a wrapper around a python dictionary. The values are the positive roots, so given a reflection, you can look up the corresponding root. If you want a list of all reflections, use the method keys for the family of reflections:

```
sage: W = WeylGroup("B3",prefix="s")
sage: [s1,s2,s3] = W.simple_reflections()
sage: ref = W.reflections(); ref
Finite family {s1*s2*s1: (1, 0, -1), s2: (0, 1, -1), s3*s2*s3: (0, 1, 1),
s3*s1*s2*s3*s1: (1, 0, 1), s1: (1, -1, 0), s2*s3*s1*s2*s3*s1*s2: (1, 1, 0),
s1*s2*s3*s2*s1: (1, 0, 0), s2*s3*s2: (0, 1, 0), s3: (0, 0, 1)}
sage: ref[s3*s2*s3]
(0, 1, 1)
sage: ref.keys()
[s1*s2*s1, s2, s3*s2*s3, s2*s3*s1*s2*s3*s1*s2, s1, s3*s1*s2*s3*s1, s1*s2*s3*s2*s1, s2*s3*s2, s3]
```

If instead you want a dictionary whose keys are the roots and whose values are the reflections, you may use the inverse family:

The Weyl group is implemented as a GAP matrix group. You therefore can display its character table. The character table is returned as a string, which you can print:

```
X.5
      3 -1 -1 . 1 3 -1 -1 -1 1 . -1
X.6
      3 1 -1 . -1 -1 3 1 -1 1 . -1
      3 1 -1 . -1 3 -1 1 1 -1
                            . -1
X.7
      3 -1 3 . -1 -1 -1 -1
                          1
X.8
                       1
      3 1 3 .
               1 -1 -1 1 -1 -1
X.9
X.10
      4 -2 . -1
               . . . 2
      4 2 . -1 . . . -2
X.11
     6 . -2 . . -2 -2 . . . . 2 6
X.12
      X.13
```

Affine Weyl groups

Affine Weyl groups may be created the same way. You simply begin with an affine Cartan type:

```
sage: W = WeylGroup(['A',2,1],prefix="s")
sage: W.cardinality()
+Infinity
sage: [s0,s1,s2] = W.simple_reflections()
sage: s0*s1*s2*s1*s0
s0*s1*s2*s1*s0
```

The affine Weyl group differs from a classical Weyl group since it is infinite. The associated classical Weyl group is a subgroup that may be extracted as follows:

```
sage: W = WeylGroup(['A',2,1],prefix="s")
sage: W1 = W.classical(); W1
Parabolic Subgroup of the Weyl Group of type ['A', 2, 1] (as a matrix group acting on the root space)
sage: W1.simple_reflections()
Finite family {1: s1, 2: s2}
```

Although W1 in this example is isomorphic to WeylGroup ("A2") it has a different matrix realization:

```
sage: for s in WeylGroup(['A', 2, 1]).classical().simple_reflections():
....: print s
....: print
[ 1 0 0]
[ 1 -1 1]
[ 0 0 1]
[ 1 0 01
[ 0 1 0]
[1 1 1 -1]
sage: for s in WeylGroup(['A',2]).simple_reflections():
. . . . :
      print s
. . . . :
        print
. . .
[0 1 0]
[1 0 0]
[0 0 1]
[1 0 0]
[0 0 1]
[0 1 0]
```

Bruhat order

The Bruhat partial order on the Weyl group may be defined as follows.

If $u, v \in W$, find a reduced expression of v into a product of simple reflections: $v = s_1 \cdots s_n$. (It is not assumed that the s_i are distinct.) If omitting some of the s_i gives a product that represents u, then $u \le v$.

The Bruhat order is implemented in Sage as a method of Coxeter groups, and so it is available for Weyl groups, classical or affine.

If $u, v \in W$ then u.bruhat_le(v) returns True if $u \leq v$ in the Bruhat order.

If $u \le v$ then the *Bruhat interval* [u, v] is defined to be the set of all t such that $u \le t \le v$. One might try to implement this as follows:

```
sage: W = WeylGroup("A2",prefix="s")
sage: [s1,s2] = W.simple_reflections()
sage: def bi(u,v) : return [t for t in W if u.bruhat_le(t) and t.bruhat_le(v)]
...
sage: bi(s1,s1*s2*s1)
[s1*s2*s1, s1*s2, s1, s2*s1]
```

This would not be a good definition since it would fail if W is affine and be inefficient of W is large. Sage has a Bruhat interval method:

```
sage: W = WeylGroup("A2",prefix="s")
sage: [s1,s2] = W.simple_reflections()
sage: W.bruhat_interval(s1,s1*s2*s1)
[s1*s2*s1, s2*s1, s1*s2, s1]
```

This works even for affine Weyl groups.

The Bruhat graph

References:

- [Carrell1994]
- [Deodhar1977]
- [Dyer1993]
- [BumpNakasuji2010]

The Bruhat graph is a structure on the Bruhat interval. Suppose that $u \le v$. The vertices of the graph are x with $u \le x \le v$. There is a vertex connecting $x, y \in [x, y]$ if $x = y \cdot r$ where r is a reflection. If this is true then either x < y or y < x.

If W is a classical Weyl group the Bruhat graph is implemented in Sage:

```
sage: W = WeylGroup("A3",prefix="s")
sage: [s1,s2,s3] = W.simple_reflections()
sage: bg = W.bruhat_graph(s2,s2*s1*s3*s2); bg
Digraph on 10 vertices
sage: bg.show3d()
```

The Bruhat graph has interesting regularity properties that were investigated by Carrell and Peterson. It is a regular graph if both the Kazhdan Lusztig polynomials $P_{u,v}$ and P_{w_0v,w_0u} are 1, where w_0 is the long Weyl group element. It is closely related to the *Deodhar conjecture*, which was proved by Deodhar, Carrell and Peterson, Dyer and Polo.

Deodhar proved that if u < v then the Bruhat interval [u,v] contains as many elements of odd length as it does of even length. We observe that often this can be strengthened: If there exists a reflection r such that left (or right) multiplication by r takes the Bruhat interval [u,v] to itself, then this gives an explicit bijection between the elements of odd and even length in [u,v].

Let us search for such reflections. Put the following commands in a file and load the file:

```
W = WeylGroup("A3",prefix="s")
[s1,s2,s3] = W.simple_reflections()
ref = W.reflections().keys()

def find_reflection(u,v):
    bi = W.bruhat_interval(u,v)
    ret = []
    for r in ref:
        if all( r*x in bi for x in bi):
            ret.append(r)
    return ret

for v in W:
    for u in W.bruhat_interval(1,v):
        if u != v:
            print u,v,find_reflection(u,v)
```

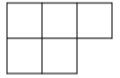
This shows that the Bruhat interval is stabilized by a reflection for all pairs (u,v) with u < v except the following two: $s_3s_1, s_1s_2s_3s_2s_1$ and $s_2, s_2s_3s_1s_2$. Now these are precisely the pairs such that $u \prec v$ in the notation of Kazhdan and Lusztig, and l(v) - l(u) > 1. One should not rashly suppose that this is a general characterization of the pairs (u,v) such that no reflection stabilizes the Bruhat interval, for this is not true. However it does suggest that the question is worthy of further investigation.

Classical Crystals

A classical crystal is one coming from the finite (classical) types $A_r, B_r, C_r, D_r, E_{6,7,8}, F_4$, and G_2 . Here we describe some background before going into the general theory of crystals and the type dependent combinatorics.

Tableaux and representations of GL(n)

Let λ be a partition. The *Young diagram* of λ is the array of boxes having λ_i boxes in the *i*-th row, left adjusted. Thus if $\lambda = (3, 2)$ the diagram is:



A semi-standard Young tableau of shape λ is a filling of the box by integers in which the rows are weakly increasing and the columns are strictly increasing. Thus

1	2	2
2	3	

is a semistandard Young tableau. Sage has a Tableau class, and you may create this tableau as follows:

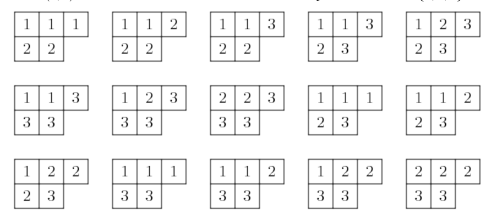
```
sage: T = Tableau([[1,2,2], [2,3]]); T
[[1, 2, 2], [2, 3]]
```

A partition of length $\leq r+1$ is a dominant weight for $GL(r+1, \mathbf{C})$ according to the description of the ambient space in Standard realizations of the ambient spaces. Therefore it corresponds to an irreducible representation $\pi_{\lambda} = \pi_{\lambda}^{GL(r+1)}$ of $GL(r+1, \mathbf{C})$.

It is true that not every dominant weight λ is a partition, since a dominant weight might have some values λ_i negative. The dominant weight λ is a partition if and only if the character of λ is a polynomial as a function on the space $\mathrm{Mat}_n(\mathbf{C})$. Thus for example $\det^{-1} = \pi_\lambda$ with $\lambda = (-1, \dots, -1)$, which is a dominant weight but not a partition, and the character is not a polynomial function on $\mathrm{Mat}_n(\mathbf{C})$.

Theorem [Littlewood] If λ is a partition, then the number of semi-standard Young tableaux with shape λ and entries in $\{1, 2, \dots, r+1\}$ is the dimension of π_{λ} .

For example, if $\lambda = (3, 2)$ and r = 2, then we find 15 tableaux with shape λ and entries in $\{1, 2, 3\}$:



This is consistent with the theorem since the dimension of the irreducible representation of GL(3) with highest weight (3, 2, 0) has dimension 15:

```
sage: A2 = WeylCharacterRing("A2")
sage: A2(3,2,0).degree()
15
```

In fact we may obtain the character of the representation from the set of tableaux. Indeed, one of the definitions of the Schur polynomial (due to Littlewood) is the following combinatorial one. If T is a tableaux, define the *weight* of T to be $\operatorname{wt}(T) = (k_1, \ldots, k_n)$ where k_i is the number of i's in the tableaux. Then the multiplicity of μ in the character χ_{λ} is the number of tableaux of weight λ . Thus if $\mathbf{z} = (z_1, \ldots, z_n)$, we have

$$\chi_{\lambda}(\mathbf{z}) = \sum_{T} \mathbf{z}^{\text{Wt}(T)}$$

where the sum is over all semi-standard Young tableaux of shape λ that have entries in $\{1, 2, \dots, r+1\}$.

Frobenius-Schur Duality

Frobenius-Schur duality is a relationship between the representation theories of the symmetric group and general linear group. We will relate this to tableaux in the next section.

Representations of the symmetric group S_k are parametrized by partitions λ of k. The parametrization may be characterized as follows. Let n be any integer $\geq k$. Then both $GL(n, \mathbf{C})$ and S_k act on $\otimes^k V$ where $V = \mathbf{C}^n$. Indeed, GL(n) acts on each V and S_k permutes them. Then if $\pi_{\lambda}^{GL(n)}$ is the representation of $GL(n, \mathbf{C})$ with highest weight

vector λ and $\pi_{\lambda}^{S_k}$ is the irreducible representation of S_k parametrized by λ then

$$\otimes^k V \cong \bigoplus_{\lambda \vdash k} \pi^{GL(n)}_{\lambda} \otimes \pi^{S_k}_{\lambda}$$

as bimodules for the two groups. This is *Frobenius-Schur duality* and it serves to characterize the parametrization of the irreducible representations of S_k by partitions of k.

Counting pairs of tableaux

In both the representation theory of GL(n) and the representation theory of S_k , the degrees of irreducible representations can be expressed in terms of the number of tableaux of the appropriate type. We have already stated the theorem for GL(n). For S_k , it goes as follows.

Let us say that a semistandard Young tableau T of shape $\lambda \vdash k$ is *standard* if T contains each entry $1, 2, \ldots, k$ exactly once. Thus both rows and columns are strictly increasing.

Theorem [Young, 1927] The degree of π_{λ} is the number of standard tableaux of shape λ .

Now let us consider the implications of Frobenius-Schur duality. The dimension of $\otimes^k V$ is n^k . Therefore n^k is equal to the number of pairs (T1, T2) of tableaux of the same shape $\lambda \vdash k$, where the first tableaux is standard (in the alphabet $1, 2, \ldots, k$), and the second the second semistandard (in the alphabet $1, 2, \ldots, n$).

The Robinson-Schensted-Knuth correspondence

The last purely combinatorial statement has a combinatorial proof, based on the Robinson-Schensted-Knuth (RSK) correspondence.

References:

- [Knuth1998], section "Tableaux and Involutions".
- [Knuth1970]
- [Fulton1997]
- [Stanley1999]

The RSK correspondence gives bijections between pairs of tableaux of various types and combinatorial objects of different types. We will not review the correspondence in detail here, but see the references. We note that Schensted insertion is implemented as the method schensted insertion of Tableau class in Sage.

Thus we have the following bijections:

- Pairs of tableaux T_1 and T_2 of shape λ where λ runs through the partitions of k such that T_1 is a standard tableau and T_2 is a semistandard tableau in $1, 2, \ldots, n$ are in bijection with the n^k words of length k in $1, 2, \ldots, n$.
- Pairs of standard tableaux of the same shape λ as λ runs through the partitions of k are in bijection with the k! elements of S_k .
- Pairs of tableaux T_1 and T_2 of the same shape λ but arbitrary size in $1, 2, 3, \ldots, n$ are in bijection with $n \times n$ positive integer matrices.
- Pairs of tableaux T_1 and T_2 of conjugate shapes λ and λ' are in bijection with $n \times n$ matrices with entries 0 or 1.

The second of these four bijection gives a combinatorial proof of the fact explained above, that the number of pairs (T1,T2) of tableaux of the same shape $\lambda \vdash k$, where the first tableaux is standard (in the alphabet $1,2,\cdots,k$), and the second the second semistandard (in the alphabet $1,2,\cdots,n$). So this second bijection is a *combinatorial analog* of Frobenius-Schur duality.

Analogies between representation theory and combinatorics

The four combinatorial bijections (variants of RSK) cited above have the following analogs in representation theory.

- The first combinatorial fact corresponds to Frobenius-Schur duality, as we have already explained.
- The second combinatorial fact also has an analog in representation theory. The group algebra $\mathbb{C}[S_k]$ is an $S_k \times S_k$ bimodule with of dimension k!. It decomposes as a direct sum of $\pi_{\lambda}^{S_k} \otimes \pi_{\lambda}^{S_k}$.

Both the combinatorial fact and the decomposition of $C[S_k]$ show that the number of pairs of standard tableaux of size k and the same shape equals k!.

• The third combinatorial fact is analogous to the decomposition of the ring of polynomial functions on $\operatorname{Mat}(n, \mathbf{C})$ on which $GL(n, \mathbf{C}) \times GL(n, \mathbf{C})$ acts by $(g_1, g_2)f(X) = f({}^tg_1Xg_2)$. The polynomial ring decomposes into the direct sum of $\pi_{\lambda}^{GL(n)} \otimes \pi_{\lambda}^{GL(n)}$.

Taking traces gives the Cauchy identity:

$$\sum_{\lambda} s_{\lambda}(x_1, \dots, x_n) s_{\lambda}(y_1, \dots, y_n) = \prod_{i,j} (1 - x_i y_j)^{-1}$$

where x_i are the eigenvalues of g_1 and y_j are the eigenvalues of g_2 . The sum is over all partitions λ .

• The last combinatorial fact is analogous to the decomposition of the exterior algebra over $Mat(n, \mathbb{C})$.

Taking traces gives the dual Cauchy identity:

$$\sum_{\lambda} s_{\lambda}(x_1, \cdots, x_n) s_{\lambda'}(y_1, \cdots, y_n) = \prod_{i,j} (1 + x_i y_j).$$

Again the sum is over partitions λ and here λ' is the conjugate partition.

Interpolating between representation theory and combinatorics

The theory of quantum groups interpolates between the representation theoretic picture and the combinatorial picture, and thereby explains these analogies. The representation $\pi_{\lambda}^{GL(n)}$ is reinterpreted as a module for the quantized enveloping algebra $U_q(\mathfrak{gl}_n(\mathbf{C}))$, and the representation $\pi_{\lambda}^{S_k}$ is reinterpreted as a module for the Iwahori Hecke algebra. Then Frobenius-Schur duality persists. See [Jimbo1986]. When $q \to 1$, we recover the representation story. When $q \to 0$, we recover the combinatorial story.

Kashiwara crystals

References:

- [Kashiwara1995]
- [KashiwaraNakashima1994]
- [HongKang2002]

Kashiwara considered the highest weight modules of quantized enveloping algebras $U_q(\mathfrak{g})$ in the limit when $q \to 0$. The enveloping algebra cannot be defined when q = 0, but a limiting structure can still be detected. This is the *crystal basis* of the module.

Kashiwara's crystal bases have a combinatorial structure that sheds light even on purely combinatorial constructions on tableaux that predated quantum groups. It gives a good generalization to other Cartan types (or more generally to Kac-Moody algebras).

Let Λ be the weight lattice of a Cartan type with root system Φ . We now define a *crystal* of type Φ . Let \mathcal{B} be a set, and let $0 \notin \mathcal{B}$ be an auxiliary element. For each index $1 \le i \le r$ we assume there given maps $e_i, f_i : \mathcal{B} \longrightarrow \mathcal{B} \cup \{0\}$, maps $\varepsilon_i, \varphi_i : \mathcal{B} \longrightarrow \mathbf{Z}$ and a map wt $: \mathcal{B} \longrightarrow \Lambda$ satisfying certain assumptions, which we now describe. It is assumed that if $x, y \in \mathcal{B}$ then $e_i(x) = y$ if and only if $f_i(y) = x$. In this case, it is assumed that

$$\operatorname{wt}(y) = \operatorname{wt}(x) + \alpha_i, \qquad \varepsilon_i(x) = \varepsilon_i(y) + 1, \qquad \varphi_i(x) = \varphi_i(y) - 1.$$

Moreover, we assume that

$$\varphi_i(x) - \varepsilon_i(x) = \langle \operatorname{wt}(x), \alpha_i^{\vee} \rangle$$

for all $x \in \mathcal{B}$.

We call a crystal regular if it satisfies the additional assumption that $\varepsilon_i(v)$ is the number of times that e_i may be applied to v, and that $\phi_i(v)$ is the number of times that f_i may be applied. That is, $\varphi_i(x) = \max\{k|f_i^k x \neq 0\}$ and $\varepsilon_i(x) = \max\{k|e_i^k(x) \neq 0\}$. Kashiwara also allows ε_i and φ_i to take the value $-\infty$.

Note: Most of the crystals that we are concerned with here are regular.

Given the crystal \mathcal{B} , the *character* $\chi_{\mathcal{B}}$ is:

$$\sum_{v \in \mathcal{B}} \mathbf{z}^{wt(v)}.$$

Given any highest weight λ , constructions of Kashiwara and Nakashima, Littelmann and others produce a crystal $\chi_{\mathcal{B}_{\lambda}}$ such that $\chi_{\mathcal{B}_{\lambda}} = \chi_{\lambda}$, where χ_{λ} is the irreducible character with highest weight λ , as in *Representations and characters*.

The crystal \mathcal{B}_{λ} is not uniquely characterized by the properties that we have stated so far. For Cartan types A, D, E (more generally, any simply-laced type) it may be characterized by these properties together with certain other *Stembridge axioms*. We will take it for granted that there is a unique "correct" crystal \mathcal{B}_{λ} and discuss how these are constructed in Sage.

Installing dot2tex

Before giving examples of crystals, we digress to help you install dot2tex, which you will need in order to make latex images of crystals.

dot2tex is an optional package of sage and the latest version can be installed via:

sage -i dot2tex

Crystals of tableaux in Sage

All crystals that are currently in Sage can be accessed by crystals. <tab>.

For type A_r , Kashiwara and Nakashima put a crystal structure on the set of tableaux with shape λ in $1, 2, \ldots, r+1$, and this is a realization of \mathcal{B}_{λ} . Moreover, this construction extends to other Cartan types, as we will explain. At the moment, we will consider how to draw pictures of these crystals.

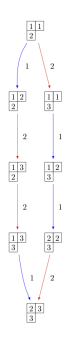
Once you have dot2tex installed, you may make images pictures of crystals with a command such as this:

```
sage: crystals.Tableaux("A2", shape=[2,1]).latex_file("/tmp/a2rho.tex") # optional - dot2tex graphvi.
```

Here $\lambda=(2,1)=(2,1,0)$. The crystal $\mathbb C$ is $\mathcal B_\lambda$. The character χ_λ will therefore be the eight-dimensional irreducible character with this highest weight. Then you may run pdflatex on the file a2rho.tex. This can also be achieved without the detour of saving the latex file via:

```
sage: B = crystals.Tableaux(['A',2], shape=[2,1])
sage: view(B, tightpage=True) # optional - dot2tex graphviz, not tested (opens external window)
```

This produces the crystal graph:



You may also wish to color the edges in different colors by specifying further latex options:

```
sage: B = crystals.Tableaux(['A',2], shape=[2,1])
sage: G = B.digraph()
sage: G.set_latex_options(color_by_label = {1:"red", 2:"yellow"})
sage: view(G, tightpage=True) # optional - dot2tex graphviz, not tested (opens external window)
```

As you can see, the elements of this crystal are exactly the eight tableaux of shape λ with entries in $\{1,2,3\}$. The convention is that if $x,y \in \mathcal{B}$ and $f_i(x)=y$, or equivalently $e_i(y)=x$, then we draw an arrow from $x\to y$. Thus the highest weight tableau is the one with no incoming arrows. Indeed, this is:

1	1
2	

We recall that the weight of the tableau is (k_1, k_2, k_3) where k_i is the number of i's in the tableau, so this tableau has weight (2, 1, 0), which indeed equals λ .

Once the crystal is created, you have access to the ambient space and its methods through the method weight_lattice_realization():

```
sage: C = crystals.Tableaux("A2", shape=[2,1])
sage: L = C.weight_lattice_realization(); L
Ambient space of the Root system of type ['A', 2]
sage: L.fundamental_weights()
Finite family {1: (1, 0, 0), 2: (1, 1, 0)}
```

The highest weight vector is available as follows:

```
sage: C = crystals.Tableaux("A2", shape=[2,1])
sage: v = C.highest_weight_vector(); v
[[1, 1], [2]]

or more simply:
sage: C = crystals.Tableaux("A2", shape=[2,1])
sage: C[0]
[[1, 1], [2]]
```

Now we may apply the operators e_i and f_i to move around in the crystal:

```
sage: C = crystals.Tableaux("A2", shape=[2,1])
sage: v = C.highest_weight_vector()
sage: v.f(1)
[[1, 2], [2]]
sage: v.f(1).f(1)
sage: v.f(1).f(1) is None
True
sage: v.f(1).f(2)
[[1, 3], [2]]
sage: v.f(1).f(2).f(2)
[[1, 3], [3]]
sage: v.f(1).f(2).f(2).f(1)
[[2, 3], [3]]
sage: v.f(1).f(2).f(2).f(1) == v.f(2).f(1).f(2).f(2)
True
```

You can construct the character if you first make a Weyl character ring:

```
sage: A2 = WeylCharacterRing("A2")
sage: C = crystals.Tableaux("A2", shape=[2,1])
sage: C.character(A2)
A2(2,1,0)
```

Crystals of letters

For each of the classical Cartan types there is a *standard crystal* $\mathcal{B}_{standard}$ from which other crystals can be built up by taking tensor products and extracting constituent irreducible crystals. This procedure is sufficient for Cartan types A_r and C_r . For types B_r and D_r the standard crystal must be supplemented with *spin crystals*. See [KashiwaraNakashima1994] or [HongKang2002] for further details.

Here is the standard crystal of type A_r .

You may create the crystal and work with it as follows:

```
sage: C = crystals.Letters("A6")
sage: v0 = C.highest_weight_vector(); v0
1
sage: v0.f(1)
2
sage: v0.f(1).f(2)
3
sage: [v0.f(1).f(2).f(x) for x in [1..6]]
```

```
[None, None, 4, None, None, None] 

sage: [v0.f(1).f(2).e(x) for x in [1..6]]

[None, 2, None, None, None, None]
```

Here is the standard crystal of type B_r .

There is, additionally, a spin crystal for B_r , corresponding to the 2^r -dimensional spin representation. We will not draw it, but we will describe it. Its elements are vectors $\epsilon_1 \otimes \cdots \otimes \epsilon_r$, where each spin $\epsilon_i = \pm$.

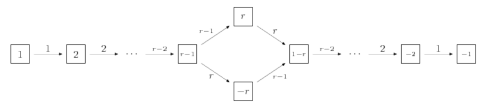
If i < r, then the effect of the operator f_i is to annihilate $v = \epsilon_1 \otimes \cdots \otimes \epsilon_r$ unless $\epsilon_i \otimes \epsilon_{i+1} = + \otimes -$. If this is so, then $f_i(v)$ is obtained from v by replacing $\epsilon_i \otimes \epsilon_{i+1}$ by $-\otimes +$. If i = r, then f_r annihilates v unless $\epsilon_r = +$, in which case it replaces ϵ_r by -.

Create the spin crystal as follows. The crystal elements are represented in the signature representation listing the ϵ_i :

```
sage: C = crystals.Spins("B3")
sage: C.list()
[+++, ++-, +-+, -++, +--, -+-, ---]
```

Here is the standard crystal of type C_r .

Here is the standard crystal of type D_r .



There are two spin crystals for type D_r . Each consists of $\epsilon_1 \otimes \cdots \otimes \epsilon_r$ with $\epsilon_i = \pm$, and the number of spins either always even or always odd. We will not describe the effect of the root operators f_i , but you are invited to create them and play around with them to guess the rule:

```
sage: Cplus = crystals.SpinsPlus("D4")
sage: Cminus = crystals.SpinsMinus("D4")
```

It is also possible to construct the standard crystal for type G_2 , E_6 , and E_7 . Here is the one for type G_2 (corresponding to the representation of degree 7):

The crystal of letters is a special case of the crystal of tableaux in the sense that $\mathcal{B}_{standard}$ is isomorphic to the crystal of tableaux whose highest weight λ is the highest weight vector of the standard representation. Thus compare:

```
sage: crystals.Letters("A3")
The crystal of letters for type ['A', 3]
sage: crystals.Tableaux("A3", shape=[1])
The crystal of tableaux of type ['A', 3] and shape(s) [[1]]
```

These two crystals are different in implementation, but they are isomorphic. In fact the second crystal is constructed from the first. We can test isomorphisms between crystals as follows:

```
sage: Cletter = crystals.Letters(['A',3])
sage: Ctableaux = crystals.Tableaux(['A',3], shape = [1])
sage: Cletter.digraph().is_isomorphic(Ctableaux.digraph())
True
sage: Cletter.digraph().is_isomorphic(Ctableaux.digraph(), certify = True)
(True, {1: [[1]], 2: [[2]], 3: [[3]], 4: [[4]]})
```

where in the last step the explicit map between the vertices of the crystals is given.

Crystals of letters have a special role in the theory since they are particularly simple, yet as Kashiwara and Nakashima showed, the crystals of tableaux can be created from them. We will review how this works.

Tensor products of crystals

Kashiwara defined the tensor product of crystals in a purely combinatorial way. The beauty of this construction is that it exactly parallels the tensor product of crystals of representations. That is, if λ and μ are dominant weights, then $\mathcal{B}_{\lambda} \otimes \mathcal{B}_{\mu}$ is a (usually disconnected) crystal, which may contain multiple copies of \mathcal{B}_{ν} (for another dominant weight ν), and the number of copies of \mathcal{B}_{ν} is exactly the multiplicity of χ_{ν} in $\chi_{\lambda}\chi_{\mu}$.

We will describe two conventions for the tensor product of crystals.

Kashiwara's definition As a set, the tensor product $\mathcal{B} \otimes \mathcal{C}$ of crystals \mathcal{B} and \mathcal{C} is the Cartesian product, but we denote the ordered pair (x,y) with $x \in \mathcal{B}$ and $y \in \mathcal{C}$ by $x \otimes y$. We define $\operatorname{wt}(x \otimes y) = \operatorname{wt}(x) + \operatorname{wt}(y)$. We define

$$f_i(x \otimes y) = \begin{cases} f_i(x) \otimes y & \text{if } \varphi_i(x) > \varepsilon_i(y), \\ x \otimes f_i(y) & \text{if } \varphi_i(x) \le \varepsilon_i(y), \end{cases}$$

and

$$e_i(x \otimes y) = \begin{cases} e_i(x) \otimes y & \text{if } \varphi_i(x) \ge \varepsilon_i(y), \\ x \otimes e_i(y) & \text{if } \varphi_i(x) < \varepsilon_i(y). \end{cases}$$

It is understood that $x \otimes 0 = 0 \otimes x = 0$. We also define:

$$\varphi_i(x \otimes y) = \max(\varphi_i(y), \varphi_i(x) + \varphi_i(y) - \varepsilon_i(y)),$$

$$\varepsilon_i(x \otimes y) = \max(\varepsilon_i(x), \varepsilon_i(x) + \varepsilon_i(y) - \varphi_i(x)).$$

Alternative definition As a set, the tensor product $\mathcal{B} \otimes \mathcal{C}$ of crystals \mathcal{B} and \mathcal{C} is the Cartesian product, but we denote the ordered pair (y, x) with $y \in \mathcal{B}$ and $x \in \mathcal{C}$ by $x \otimes y$. We define $\operatorname{wt}(x \otimes y) = \operatorname{wt}(y) + \operatorname{wt}(x)$. We define

$$f_i(x \otimes y) = \begin{cases} f_i(x) \otimes y & \text{if } \varphi_i(y) \leq \varepsilon_i(x), \\ x \otimes f_i(y) & \text{if } \varphi_i(y) > \varepsilon_i(x), \end{cases}$$

and

$$e_i(x \otimes y) = \begin{cases} e_i(x) \otimes y & \text{if } \varphi_i(y) < \varepsilon_i(x), \\ x \otimes e_i(y) & \text{if } \varphi_i(y) \ge \varepsilon_i(x). \end{cases}$$

It is understood that $y \otimes 0 = 0 \otimes y = 0$. We also define

$$\varphi_i(x \otimes y) = \max(\varphi_i(x), \varphi_i(y) + \varphi_i(x) - \varepsilon_i(x)),$$

$$\varepsilon_i(x \otimes y) = \max(\varepsilon_i(y), \varepsilon_i(y) + \varepsilon_i(x) - \varphi_i(y)).$$

The tensor product is associative: $(x \otimes y) \otimes z \mapsto x \otimes (y \otimes z)$ is an isomorphism $(\mathcal{B} \otimes \mathcal{C}) \otimes \mathcal{D} \to \mathcal{B} \otimes (\mathcal{C} \otimes \mathcal{D})$, and so we may consider tensor products of arbitrary numbers of crystals.

The relationship between the two definitions The relationship between the two definitions is simply that the Kashiwara tensor product $\mathcal{B} \otimes \mathcal{C}$ is the alternate tensor product $\mathcal{C} \otimes \mathcal{B}$ in reverse order. Sage uses the alternative tensor product. Even though the tensor product construction is *a priori* asymmetrical, both constructions produce isomorphic crystals, and in particular Sage's crystals of tableaux are identical to Kashiwara's.

Note: Using abstract crystals (i.e. they satisfy the axioms but do not arise from a representation of $U_q(\mathfrak{g})$), we can construct crystals \mathcal{B}, \mathcal{C} such that $\mathcal{B} \otimes \mathcal{C} \neq \mathcal{C} \otimes \mathcal{B}$ (of course, using the same convention).

Tensor products of crystals in Sage You may construct the tensor product of several crystals in Sage using crystals. TensorProduct:

```
sage: C = crystals.Letters("A2")
sage: T = crystals.TensorProduct(C,C,C); T
Full tensor product of the crystals [The crystal of letters for type ['A', 2],
The crystal of letters for type ['A', 2], The crystal of letters for type ['A', 2]]
sage: T.cardinality()
27
sage: T.highest_weight_vectors()
([1, 1, 1], [2, 1, 1], [1, 2, 1], [3, 2, 1])
```

This crystal has four highest weight vectors. We may understand this as follows:

```
sage: A2 = WeylCharacterRing("A2")
sage: C = crystals.Letters("A2")
sage: T = crystals.TensorProduct(C,C,C)
sage: chi_C = C.character(A2)
sage: chi_T = T.character(A2)
sage: chi_C
A2(1,0,0)
sage: chi_T
A2(1,1,1) + 2*A2(2,1,0) + A2(3,0,0)
sage: chi_T == chi_C^3
True
```

As expected, the character of T is the cube of the character of C, and representations with highest weight (1,1,1), (3,0,0) and (2,1,0). This decomposition is predicted by Frobenius-Schur duality: the multiplicity of $\pi_{\lambda}^{GL(n)}$ in $\otimes^3 \mathbf{C}^3$ is the degree of $\pi_{\lambda}^{S_3}$.

It is useful to be able to select one irreducible constitutent of T. If we only want one of the irreducible constituents of T, we can specify a list of highest weight vectors by the option generators. If the list has only one element, then we get an irreducible crystal. We can make four such crystals:

```
sage: A2 = WeylCharacterRing("A2")
sage: C = crystals.Letters("A2")
sage: T = crystals.TensorProduct(C,C,C)
sage: [T1,T2,T3,T4] = \
    [crystals.TensorProduct(C,C,C,generators=[v]) for v in T.highest_weight_vectors()]
sage: [B.cardinality() for B in [T1,T2,T3,T4]]
[10, 8, 8, 1]
sage: [B.character(A2) for B in [T1,T2,T3,T4]]
[A2(3,0,0), A2(2,1,0), A2(2,1,0), A2(1,1,1)]
```

We see that two of these crystals are isomorphic, with character A2 (2, 1, 0). Try:

```
sage: A2 = WeylCharacterRing("A2")
sage: C = crystals.Letters("A2")
```

```
sage: T = crystals.TensorProduct(C,C,C)
sage: [T1,T2,T3,T4] = \
    [crystals.TensorProduct(C,C,C,generators=[v]) for v in T.highest_weight_vectors()]
sage: T1.plot()
Graphics object consisting of 35 graphics primitives
sage: T2.plot()
Graphics object consisting of 25 graphics primitives
sage: T3.plot()
Graphics object consisting of 25 graphics primitives
sage: T4.plot()
Graphics object consisting of 2 graphics primitives
```

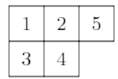
Elements of crystals. TensorProduct (A, B, C, ...) are represented by sequences [a, b, c, ...] with a in A, b in B, etc. This of course represents $a \otimes b \otimes c \otimes \cdots$.

Crystals of tableaux as tensor products of crystals

Sage implements the CrystalOfTableaux as a subcrystal of a tensor product of the ClassicalCrystalOfLetters. You can see how its done as follows:

```
sage: T = crystals.Tableaux("A4", shape=[3,2])
sage: v = T.highest_weight_vector().f(1).f(2).f(3).f(2).f(1).f(4).f(2).f(3); v
[[1, 2, 5], [3, 4]]
sage: v._list
[3, 1, 4, 2, 5]
```

We've looked at the internal representation of v, where it is represented as an element of the fourth tensor power of the ClassicalCrystalOfLetters. We see that the tableau:



is interpreted as the tensor:



The elements of the tableau are read from bottom to top and from left to right. This is the *inverse middle-Eastern* reading of the tableau. See Hong and Kang, *loc. cit.* for discussion of the readings of a tableau.

Spin crystals

For the Cartan types A_r , C_r or G_2 , CrystalOfTableaux are capable of making any finite crystal. (For type A_r it is necessary that the highest weight λ be a partition.)

For Cartan types B_r and D_r , there also exist spin representations. The corresponding crystals are implemented as *spin crystals*. For these types, CrystalOfTableaux also allows the input shape λ to be half-integral if it is of height r. For example:

```
sage: C = crystals.Tableaux(['B',2], shape = [3/2, 1/2])
sage: C.list()
[[++, [[1]]], [++, [[2]]], [++, [[0]]], [++, [[-2]]], [++, [[-1]]], [+-, [[-2]]],
```

```
[+-, [[-1]]], [-+, [[-1]]], [+-, [[1]]], [+-, [[2]]], [-+, [[2]]], [+-, [[0]]], [-+, [[0]]], [--, [[-2]]], [--, [[-1]]]]
```

Here the first list of + and - gives a spin column that is discussed in more detail in the next section and the second entry is a crystal of tableau element for $\lambda = (\lfloor \lambda_1 \rfloor, \lfloor \lambda_2 \rfloor, \ldots)$. For type D_r , we have the additional feature that there are two types of spin crystals. Hence in CrystalOfTableaux the r-th entry of λ in this case can also take negative values:

```
sage: C = crystals.Tableaux(['D',3], shape = [1/2, 1/2,-1/2])
sage: C.list()
[[++-, []], [+-+, []], [-++, []]]
```

For rank two Cartan types, we also have crystals. FastRankTwo which gives a different fast implementation of these crystals:

```
sage: B = crystals.FastRankTwo(['B',2], shape=[3/2,1/2]); B
The fast crystal for B2 with shape [3/2,1/2]
sage: v = B.highest_weight_vector(); v.weight()
(3/2, 1/2)
```

Type B spin crystal The spin crystal has highest weight (1/2, ..., 1/2). This is the last fundamental weight. The irreducible representation with this weight is the spin representation of degree 2^r . Its crystal is hand-coded in Sage:

```
sage: Cspin = crystals.Spins("B3"); Cspin
The crystal of spins for type ['B', 3]
sage: Cspin.cardinality()
```

The crystals with highest weight λ , where λ is a half-integral weight, are constructed as a tensor product of a spin column and the highest weight crystal of the integer part of λ . For example, suppose that $\lambda = (3/2, 3/2, 1/2)$. The corresponding irreducible character will have degree 112:

```
sage: B3 = WeylCharacterRing("B3")
sage: B3(3/2,3/2,1/2).degree()
112
```

So \mathcal{B}_{λ} will have 112 elements. We can find it as a subcrystal of Cspin $\otimes \mathcal{B}_{\mu}$, where $\mu = \lambda - (1/2, 1/2, 1/2) = (1, 1, 0)$:

```
sage: B3 = WeylCharacterRing("B3")
sage: B3(1,1,0)*B3(1/2,1/2,1/2)
B3(1/2,1/2,1/2) + B3(3/2,1/2,1/2) + B3(3/2,3/2,1/2)
```

We see that just taking the tensor product of these two crystals will produce a reducible crystal with three constitutents, and we want to extract the one we want. We do that as follows:

```
sage: B3 = WeylCharacterRing("B3")
sage: C1 = crystals.Tableaux("B3", shape=[1,1])
sage: Cspin = crystals.Spins("B3")
sage: C = crystals.TensorProduct(C1, Cspin, generators=[[C1[0],Cspin[0]]])
sage: C.cardinality()
112
```

Alternatively, we can get this directly from CrystalOfTableaux:

```
sage: C = crystals.Tableaux(['B',3], shape = [3/2,3/2,1/2])
sage: C.cardinality()
112
```

This is the desired crystal.

Type D spin crystals A similar situation pertains for type D_r , but now there are two spin crystals, both of degree 2^{r-1} . These are hand-coded in sage:

```
sage: SpinPlus = crystals.SpinsPlus("D4")
sage: SpinMinus = crystals.SpinsMinus("D4")
sage: SpinPlus[0].weight()
(1/2, 1/2, 1/2, 1/2)
sage: SpinMinus[0].weight()
(1/2, 1/2, 1/2, -1/2)
sage: [C.cardinality() for C in [SpinPlus,SpinMinus]]
[8, 8]
```

Similarly to type B crystal, we obtain crystal with spin weight by allowing for partitions with half-integer values, and the last entry can be negative depending on the type of the spin.

Lusztig involution

The Lusztig involution on a finite-dimensional highest weight crystal $B(\lambda)$ of highest weight λ maps the highest weight vector to the lowest weight vector and the Kashiwara operator f_i to e_{i^*} , where i^* is defined as $\alpha_{i^*} = -w_0(\alpha_i)$. Here w_0 is the longest element of the Weyl group acting on the i-th simple root α_i . For example, for type A_n we have $i^* = n + 1 - i$, whereas for type C_n we have $i^* = i$. For type D_n and n even also have $i^* = i$, but for n odd this map interchanges nodes n - 1 and n. Here is how to achieve this in Sage:

```
sage: B = crystals.Tableaux(['A',3],shape=[2,1])
sage: b = B(rows=[[1,2],[3]])
sage: b.lusztig_involution()
[[2, 4], [3]]
```

For type A_n , the Lusztig involution is the same as the Schuetzenberger involution (which in Sage is defined on tableaux):

```
sage: t = Tableau([[1,2],[3]])
sage: t.schuetzenberger_involution(n=4)
[[2, 4], [3]]
```

For all tableaux in a given crystal, this can be tested via:

```
sage: B = crystals.Tableaux(['A',3],shape=[2])
sage: all(b.lusztig_involution().to_tableau() == b.to_tableau().schuetzenberger_involution(n=4) for large
```

The Lusztig involution is also defined for finite-dimensional highest weight crystals of exceptional type:

```
sage: C = CartanType(['E',6])
sage: La = C.root_system().weight_lattice().fundamental_weights()
sage: T = crystals.HighestWeight(La[1])
sage: t = T[4]; t
[(-2, 5)]
sage: t.lusztig_involution()
[(-3, 2)]
```

Levi branching rules for crystals

Let G be a Lie group and H a Levi subgroup. We have already seen that the Dynkin diagram of H is obtained from that of G by erasing one or more nodes.

If C is a crystal for G, then we may obtain the corresponding crystal for H by a similar process. For example if the Dynkin diagram for H is obtained from the Dynkin diagram for G by erasing the i-th node, then if we erase all the edges in the crystal C that are labeled with i, we obtain a crystal for H.

In Sage this is achieved by specifying the index set used in the digraph method:

```
sage: T = crystals.Tableaux(['D',4],shape=[1])
sage: G = T.digraph(index_set=[1,2,3])
```

We see that the type D_4 crystal indeed decomposes into two type A_3 components.



For more on branching rules, see *Maximal Subgroups and Branching Rules* or *Levi subgroups* for specifics on the Levi subgroups.

Subcrystals

Sometimes it might be desirable to work with a subcrystal of a crystal. For example, one might want to look at all $\{2, 3, ..., n\}$ highest elements of a crystal and look at a particular such component:

```
sage: T = crystals.Tableaux(['D',4],shape=[2,1])
sage: hw = [ t for t in T if t.is_highest_weight(index_set = [2,3,4]) ]; hw
[[[1, 1], [2]],
[[1, 2], [2]],
[[2, -1], [-2]],
[[2, -1], [-1]],
[[1, -1], [2]],
[[2, -1], [3]],
[[1, 2], [3]],
[[1, 2], [3]],
[[1, 2], [-2]],
[[2, 2], [-2]],
[[2, 2], [-1]]]
```

```
sage: C = T.subcrystal(generators = [T(rows=[[2,-1],[3]])], index_set = [2,3,4])
sage: G = T.digraph(subset = C, index_set=[2,3,4])
```

Affine Finite Crystals

In this document we briefly explain the construction and implementation of the Kirillov–Reshetikhin crystals of [FourierEtAl2009].

Kirillov-Reshetikhin (KR) crystals are finite-dimensional affine crystals corresponding to Kirillov-Reshetikhin modules. They were first conjectured to exist in [HatayamaEtAl2001]. The proof of their existence for nonexceptional types was given in [OkadoSchilling2008] and their combinatorial models were constructed in [FourierEtAl2009]. Kirillov-Reshetikhin crystals $B^{r,s}$ are indexed first by their type (like $A_n^{(1)}$, $B_n^{(1)}$, ...) with underlying index set $I = \{0, 1, \ldots, n\}$ and two integers r and s. The integers s only needs to satisfy s > 0, whereas r is a node of the finite Dynkin diagram $r \in I \setminus \{0\}$.

Their construction relies on several cases which we discuss separately. In all cases when removing the zero arrows, the crystal decomposes as a (direct sum of) classical crystals which gives the crystal structure for the index set $I_0 = \{1, 2, ..., n\}$. Then the zero arrows are added by either exploiting a symmetry of the Dynkin diagram or by using embeddings of crystals.

Type $A_n^{(1)}$

The Dynkin diagram for affine type A has a rotational symmetry mapping $\sigma: i \mapsto i+1$ where we view the indices modulo n+1:

The classical decomposition of $B^{r,s}$ is the A_n highest weight crystal $B(s\omega_r)$ or equivalently the crystal of tableaux labelled by the rectangular partition (s^r) :

$$B^{r,s} \cong B(s\omega_r)$$
 as a $\{1, 2, \dots, n\}$ -crystal

In Sage we can see this via:

```
sage: K = crystals.KirillovReshetikhin(['A',3,1],1,1)
sage: K.classical_decomposition()
The crystal of tableaux of type ['A', 3] and shape(s) [[1]]
sage: K.list()
[[[1]], [[2]], [[3]], [[4]]]
sage: K = crystals.KirillovReshetikhin(['A',3,1],2,1)
sage: K.classical_decomposition()
The crystal of tableaux of type ['A', 3] and shape(s) [[1, 1]]
```

One can change between the classical and affine crystal using the methods lift and retract:

```
sage: K = crystals.KirillovReshetikhin(['A',3,1],2,1)
sage: b = K(rows=[[1],[3]]); type(b)
<class 'sage.combinat.crystals.kirillov_reshetikhin.KR_type_A_with_category.element_class'>
sage: b.lift()
[[1], [3]]
sage: type(b.lift())
<class 'sage.combinat.crystals.tensor_product.CrystalOfTableaux_with_category.element_class'>
sage: b = crystals.Tableaux(['A',3], shape = [1,1]) (rows=[[1],[3]])
sage: K.retract(b)
[[1], [3]]
sage: type(K.retract(b))
<class 'sage.combinat.crystals.kirillov_reshetikhin.KR_type_A_with_category.element_class'>
```

The 0-arrows are obtained using the analogue of σ , called the promotion operator pr, on the level of crystals via:

$$f_0 = \operatorname{pr}^{-1} \circ f_1 \circ \operatorname{pr}$$

 $e_0 = \operatorname{pr}^{-1} \circ e_1 \circ \operatorname{pr}$

In Sage this can be achieved as follows:

```
sage: K = crystals.KirillovReshetikhin(['A',3,1],2,1)
sage: b = K.module_generator(); b

[[1], [2]]
sage: b.f(0)
sage: b.e(0)
[[2], [4]]

sage: K.promotion()(b.lift())
[[2], [3]]
sage: K.promotion()(b.lift()).e(1)
[[1], [3]]
sage: K.promotion_inverse()(K.promotion()(b.lift()).e(1))
[[2], [4]]
```

KR crystals are level 0 crystals, meaning that the weight of all elements in these crystals is zero:

```
sage: K = crystals.KirillovReshetikhin(['A',3,1],2,1)
sage: b = K.module_generator(); b.weight()
-Lambda[0] + Lambda[2]
sage: b.weight().level()
0
```

The KR crystal ${\cal B}^{1,1}$ of type ${\cal A}_2^{(1)}$ looks as follows:



In Sage this can be obtained via:

```
sage: K = crystals.KirillovReshetikhin(['A',2,1],1,1)
sage: G = K.digraph()
sage: view(G, tightpage=True) # optional - dot2tex graphviz, not tested (opens external window)
```

```
Types D_n^{(1)}, B_n^{(1)}, A_{2n-1}^{(2)}
```

The Dynkin diagrams for types $D_n^{(1)}$, $B_n^{(1)}$, $A_{2n-1}^{(2)}$ are invariant under interchanging nodes 0 and 1:

```
sage: n = 5
sage: C = CartanType(['D',n,1]); C.dynkin_diagram()
   0---0
   2 3 4
1
D5~
sage: C = CartanType(['B',n,1]); C.dynkin_diagram()
   0---0--0=>=0
1 2 3 4 5
B5~
sage: C = CartanType(['A',2*n-1,2]); C.dynkin_diagram()
0---0--0=<=0
1 2 3 4
B5~*
```

The underlying classical algebras obtained when removing node 0 are type $\mathfrak{g}_0 = D_n, B_n, C_n$, respectively. The classical decomposition into a \mathfrak{g}_0 crystal is a direct sum:

$$B^{r,s} \cong \bigoplus_{\lambda} B(\lambda)$$
 as a $\{1,2,\ldots,n\}$ -crystal

where λ is obtained from $s\omega_r$ (or equivalently a rectangular partition of shape (s^r)) by removing vertical dominoes. This in fact only holds in the ranges $1 \le r \le n-2$ for type $D_n^{(1)}$, and $1 \le r \le n$ for types $B_n^{(1)}$ and $A_{2n-1}^{(2)}$:

```
sage: K = crystals.KirillovReshetikhin(['D',6,1],4,2)
sage: K.classical_decomposition()
The crystal of tableaux of type ['D', 6] and shape(s)
[[], [1, 1], [1, 1, 1, 1], [2, 2], [2, 2, 1, 1], [2, 2, 2, 2]]
```

For type $B_n^{(1)}$ and r=n, one needs to be aware that ω_n is a spin weight and hence corresponds in the partition language to a column of height n and width 1/2:

```
sage: K = crystals.KirillovReshetikhin(['B',3,1],3,1)
sage: K.classical_decomposition()
The crystal of tableaux of type ['B', 3] and shape(s) [[1/2, 1/2, 1/2]]
```

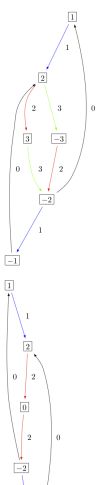
As for type $A_n^{(1)}$, the Dynkin automorphism induces a promotion-type operator σ on the level of crystals. In this case in can however happen that the automorphism changes between classical components:

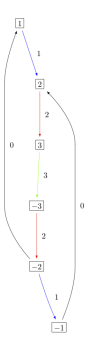
```
sage: K = crystals.KirillovReshetikhin(['D',4,1],2,1)
sage: b = K.module_generator(); b
[[1], [2]]
sage: K.automorphism(b)
[[2], [-1]]
sage: b = K(rows=[[2],[-2]])
sage: K.automorphism(b)
[]
```

This operator σ is used to define the affine crystal operators:

$$f_0 = \sigma \circ f_1 \circ \sigma$$
$$e_0 = \sigma \circ e_1 \circ \sigma$$

The KR crystals $B^{1,1}$ of types $D_3^{(1)},\, B_2^{(1)},\,$ and $A_5^{(2)}$ are, respectively:





Type $C_n^{(1)}$

The Dynkin diagram of type $C_n^{(1)}$ has a symmetry $\sigma(i) = n - i$:

```
sage: C = CartanType(['C',4,1]); C.dynkin_diagram()
0=>=0--0--0=<=0
0    1    2    3    4
C4~</pre>
```

The classical subalgebra when removing the 0 node is of type C_n .

However, in this case the crystal $B^{r,s}$ is not constructed using σ , but rather using a virtual crystal construction. $B^{r,s}$ of type $C_n^{(1)}$ is realized inside $\hat{V}^{r,s}$ of type $A_{2n+1}^{(2)}$ using:

$$e_0=\hat{e}_0\hat{e}_1$$
 and $e_i=\hat{e}_{i+1}$ for $1\leq i\leq n$ $f_0=\hat{f}_0\hat{f}_1$ and $f_i=\hat{f}_{i+1}$ for $1\leq i\leq n$

where \hat{e}_i and \hat{f}_i are the crystal operator in the ambient crystal $\hat{V}^{r,s}$:

```
sage: K = crystals.KirillovReshetikhin(['C',3,1],1,2); K.ambient_crystal()
Kirillov-Reshetikhin crystal of type ['B', 4, 1]^* with (r,s)=(1,2)
```

The classical decomposition for $1 \le r < n$ is given by:

$$B^{r,s} \cong \bigoplus_{\lambda} B(\lambda) \quad \text{as a } \{1,2,\ldots,n\}\text{-crystal}$$

where λ is obtained from $s\omega_r$ (or equivalently a rectangular partition of shape (s^r)) by removing horizontal dominoes:

```
sage: K = crystals.KirillovReshetikhin(['C',3,1],2,4)
sage: K.classical_decomposition()
The crystal of tableaux of type ['C', 3] and shape(s) [[], [2], [4], [2, 2], [4, 2], [4, 4]]
```

The KR crystal $B^{1,1}$ of type $C_2^{(1)}$ looks as follows:



Types
$$D_{n+1}^{(2)}$$
, $A_{2n}^{(2)}$

The Dynkin diagrams of types $D_{n+1}^{(2)}$ and $A_{2n}^{(2)}$ look as follows:

```
sage: C = CartanType(['D',5,2]); C.dynkin_diagram()
0=<=0---0--0=>=0
0    1    2    3    4
C4~*

sage: C = CartanType(['A',8,2]); C.dynkin_diagram()
0=<=0---0--0=<=0
0    1    2    3    4
BC4~</pre>
```

The classical subdiagram is of type B_n for type $D_{n+1}^{(2)}$ and of type C_n for type $A_{2n}^{(2)}$. The classical decomposition for these KR crystals for $1 \le r < n$ for type $D_{n+1}^{(2)}$ and $1 \le r \le n$ for type $A_{2n}^{(2)}$ is given by:

$$B^{r,s} \cong \bigoplus_{\lambda} B(\lambda)$$
 as a $\{1,2,\ldots,n\}$ -crystal

where λ is obtained from $s\omega_r$ (or equivalently a rectangular partition of shape (s^r)) by removing single boxes:

```
sage: K = crystals.KirillovReshetikhin(['D',5,2],2,2)
sage: K.classical_decomposition()
The crystal of tableaux of type ['B', 4] and shape(s) [[], [1], [2], [1, 1], [2, 1], [2, 2]]
sage: K = crystals.KirillovReshetikhin(['A',8,2],2,2)
sage: K.classical_decomposition()
The crystal of tableaux of type ['C', 4] and shape(s) [[], [1], [2], [1, 1], [2, 1], [2, 2]]
```

The KR crystals are constructed using an injective map into a KR crystal of type $C_n^{(1)}$

$$S:B^{r,s}\to B^{r,2s}_{C^{(1)}_n}\quad \text{ such that } S(e_ib)=e_i^{m_i}S(b) \text{ and } S(f_ib)=f_i^{m_i}S(b)$$

where

$$(m_0,\ldots,m_n)=(1,2,\ldots,2,1)$$
 for type $D_{n+1}^{(2)}$ and $(1,2,\ldots,2,2)$ for type $A_{2n}^{(2)}$.

```
sage: K = crystals.KirillovReshetikhin(['D',5,2],1,2); K.ambient_crystal()
Kirillov-Reshetikhin crystal of type ['C', 4, 1] with (r,s)=(1,4)
sage: K = crystals.KirillovReshetikhin(['A',8,2],1,2); K.ambient_crystal()
Kirillov-Reshetikhin crystal of type ['C', 4, 1] with (r,s)=(1,4)
```

The KR crystals $B^{1,1}$ of type $D_3^{(2)}$ and $A_4^{(2)}$ look as follows:



As you can see from the Dynkin diagram for type $A_{2n}^{(2)}$, mapping the nodes $i\mapsto n-i$ yields the same diagram, but with relabelled nodes. In this case the classical subdiagram is of type B_n instead of C_n . One can also construct the KR crystal $B^{r,s}$ of type $A_{2n}^{(2)}$ based on this classical decomposition. In this case the classical decomposition is the sum over all weights obtained from $s\omega_r$ by removing horizontal dominoes:

```
sage: C = CartanType(['A',6,2]).dual()
sage: Kdual = crystals.KirillovReshetikhin(C,2,2)
sage: Kdual.classical_decomposition()
The crystal of tableaux of type ['B', 3] and shape(s) [[], [2], [2, 2]]
```

Looking at the picture, one can see that this implementation is isomorphic to the other implementation based on the C_n decomposition up to a relabeling of the arrows:

```
sage: C = CartanType(['A', 4, 2])
sage: K = crystals.KirillovReshetikhin(C, 1, 1)
sage: Kdual = crystals.KirillovReshetikhin(C.dual(), 1, 1)
sage: G = K.digraph()
sage: Gdual = Kdual.digraph()
```

```
sage: f = { 1:1, 0:2, 2:0 }
sage: for u,v,label in Gdual.edges():
....: Gdual.set_edge_label(u,v,f[label])
sage: G.is_isomorphic(Gdual, edge_labels = True)
True
```



Exceptional nodes

The KR crystals $B^{n,s}$ for types $C_n^{(1)}$ and $D_{n+1}^{(2)}$ were excluded from the above discussion. They are associated to the exceptional node r=n and in this case the classical decomposition is irreducible:

$$B^{n,s} \cong B(s\omega_n).$$

In Sage:

```
sage: K = crystals.KirillovReshetikhin(['C',2,1],2,1)
sage: K.classical_decomposition()
The crystal of tableaux of type ['C', 2] and shape(s) [[1, 1]]
sage: K = crystals.KirillovReshetikhin(['D',3,2],2,1)
sage: K.classical_decomposition()
The crystal of tableaux of type ['B', 2] and shape(s) [[1/2, 1/2]]
```



The KR crystals $B^{n,s}$ and $B^{n-1,s}$ of type $D_n^{(1)}$ are also special. They decompose as:

$$B^{n,s} \cong B(s\omega_n)$$
 and $B^{n-1,s} \cong B(s\omega_{n-1})$.

```
sage: K = crystals.KirillovReshetikhin(['D',4,1],4,1)
sage: K.classical_decomposition()
The crystal of tableaux of type ['D', 4] and shape(s) [[1/2, 1/2, 1/2, 1/2]]
sage: K = crystals.KirillovReshetikhin(['D',4,1],3,1)
sage: K.classical_decomposition()
The crystal of tableaux of type ['D', 4] and shape(s) [[1/2, 1/2, 1/2, -1/2]]
```

Type $E_6^{(1)}$

In [JonesEtAl2010] the KR crystals $B^{r,s}$ for r=1,2,6 in type $E_6^{(1)}$ were constructed exploiting again a Dynkin diagram automorphism, namely the automorphism σ of order 3 which maps $0\mapsto 1\mapsto 6\mapsto 0$:

The crystals $B^{1,s}$ and $B^{6,s}$ are irreducible as classical crystals:

```
sage: K = crystals.KirillovReshetikhin(['E',6,1],1,1)
sage: K.classical_decomposition()
Direct sum of the crystals Family (Finite dimensional highest weight crystal of type ['E', 6] and highest K = crystals.KirillovReshetikhin(['E',6,1],6,1)
sage: K.classical_decomposition()
Direct sum of the crystals Family (Finite dimensional highest weight crystal of type ['E', 6] and highest weight crystal of type ['E', 6]
```

whereas for the adjoint node r = 2 we have the decomposition

$$B^{2,s} \cong \bigoplus_{k=0}^{s} B(k\omega_2)$$

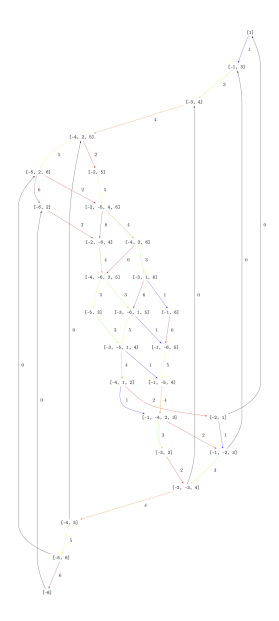
```
sage: K = crystals.KirillovReshetikhin(['E',6,1],2,1)
sage: K.classical_decomposition()
Direct sum of the crystals Family (Finite dimensional highest weight crystal of type ['E', 6] and highest weight Lambda[2])
```

The promotion operator on the crystal corresponding to σ can be calculated explicitly:

```
sage: K = crystals.KirillovReshetikhin(['E',6,1],1,1)
sage: promotion = K.promotion()
sage: u = K.module_generator(); u
[(1,)]
sage: promotion(u.lift())
[(-1, 6)]
```

The crystal $B^{1,1}$ is already of dimension 27. The elements b of this crystal are labelled by tuples which specify their nonzero $\phi_i(b)$ and $\epsilon_i(b)$. For example, [-6,2] indicates that $\phi_2([-6,2]) = \epsilon_6([-6,2]) = 1$ and all others are equal to zero:

```
sage: K = crystals.KirillovReshetikhin(['E',6,1],1,1)
sage: K.cardinality()
27
```



Single column KR crystals

A single column KR crystal is $B^{r,1}$ for any $r \in I_0$.

In [LNSSS14II] and [LNSSS14II], it was shown that single column KR crystals can be constructed by projecting level 0 crystals of LS paths onto the classical weight lattice. We first verify that we do get an isomorphic crystal for $B^{1,1}$ in type $E_6^{(1)}$:

```
sage: K = crystals.KirillovReshetikhin(['E',6,1], 1,1)
sage: K2 = crystals.kirillov_reshetikhin.LSPaths(['E',6,1], 1,1)
sage: K.digraph().is_isomorphic(K2.digraph(), edge_labels=True)
True
```

Here is an example in $E_8^{(1)}$ and we calculate its classical decomposition:

```
sage: K = crystals.kirillov_reshetikhin.LSPaths(['E',8,1], 8,1)
sage: K.cardinality()
249
```

```
sage: L = [x for x in K if x.is_highest_weight([1,2,3,4,5,6,7,8])]
sage: map(lambda x: x.weight(), L)
[-2*Lambda[0] + Lambda[8], 0]
```

Applications

An important notion for finite-dimensional affine crystals is perfectness. The crucial property is that a crystal B is perfect of level ℓ if there is a bijection between level ℓ dominant weights and elements in

$$B_{\min} = \{ b \in B \mid \text{lev}(\varphi(b)) = \ell \}$$
.

For a precise definition of perfect crystals see [HongKang2002] . In [FourierEtAl2010] it was proven that for the nonexceptional types $B^{r,s}$ is perfect as long as s/c_r is an integer. Here $c_r=1$ except $c_r=2$ for $1 \le r < n$ in type $C_n^{(1)}$ and r=n in type $B_n^{(1)}$.

Here we verify this using Sage for $B^{1,1}$ of type $C_3^{(1)}$:

```
sage: K = crystals.KirillovReshetikhin(['C',3,1],1,1)
sage: Lambda = K.weight_lattice_realization().fundamental_weights(); Lambda
Finite family {0: Lambda[0], 1: Lambda[1], 2: Lambda[2], 3: Lambda[3]}
sage: [w.level() for w in Lambda]
[1, 1, 1, 1]
sage: Bmin = [b for b in K if b.Phi().level() == 1 ]; Bmin
[[[1]], [[2]], [[3]], [[-3]], [[-2]], [[-1]]]
sage: [b.Phi() for b in Bmin]
[Lambda[1], Lambda[2], Lambda[3], Lambda[2], Lambda[1], Lambda[0]]
```

As you can see, both b=1 and b=-2 satisfy $\varphi(b)=\Lambda_1$. Hence there is no bijection between the minimal elements in B_{\min} and level 1 weights. Therefore, $B^{1,1}$ of type $C_3^{(1)}$ is not perfect. However, $B^{1,2}$ of type $C_n^{(1)}$ is a perfect crystal:

```
sage: K = crystals.KirillovReshetikhin(['C',3,1],1,2)
sage: Lambda = K.weight_lattice_realization().fundamental_weights()
sage: Bmin = [b for b in K if b.Phi().level() == 1 ]
sage: [b.Phi() for b in Bmin]
[Lambda[0], Lambda[3], Lambda[2], Lambda[1]]
```

Perfect crystals can be used to construct infinite-dimensional highest weight crystals and Demazure crystals using the Kyoto path model [KKMMNN1992]. We construct Example 10.6.5 in [HongKang2002]:

```
sage: K = crystals.KirillovReshetikhin(['A',1,1], 1,1)
sage: La = RootSystem(['A',1,1]).weight_lattice().fundamental_weights()
sage: B = crystals.KyotoPathModel(K, La[0])
sage: B.highest_weight_vector()
[[[2]]]

sage: K = crystals.KirillovReshetikhin(['A',2,1], 1,1)
sage: La = RootSystem(['A',2,1]).weight_lattice().fundamental_weights()
sage: B = crystals.KyotoPathModel(K, La[0])
sage: B.highest_weight_vector()
[[[3]]]

sage: K = crystals.KirillovReshetikhin(['C',2,1], 2,1)
sage: La = RootSystem(['C',2,1]).weight_lattice().fundamental_weights()
sage: B = crystals.KyotoPathModel(K, La[1])
```

```
sage: B.highest_weight_vector()
[[[2], [-2]]]
```

Energy function and one-dimensional configuration sum

For tensor products of Kirillov-Reshehtikhin crystals, there also exists the important notion of the energy function. It can be defined as the sum of certain local energy functions and the R-matrix. In Theorem 7.5 in [SchillingTingley2011] it was shown that for perfect crystals of the same level the energy D(b) is the same as the affine grading (up to a normalization). The affine grading is defined as the minimal number of applications of e_0 to b to reach a ground state path. Computationally, this algorithm is a lot more efficient than the computation involving the R-matrix and has been implemented in Sage:

```
sage: K = crystals.KirillovReshetikhin(['A',2,1],1,1)
sage: T = crystals.TensorProduct(K,K,K)
sage: hw = [b for b in T if all(b.epsilon(i)==0 for i in [1,2])]
sage: for b in hw:
...:     print b, b.energy_function()
[[[1]], [[1]], [[1]]] 0
[[[1]], [[2]], [[1]]] 2
[[[2]], [[1]], [[1]]] 1
[[[3]], [[2]], [[1]]] 3
```

The affine grading can be computed even for nonperfect crystals:

The one-dimensional configuration sum of a crystal B is the graded sum by energy of the weight of all elements $b \in B$:

$$X(B) = \sum_{b \in B} x^{\text{weight}(b)} q^{D(b)}$$

Here is an example of how you can compute the one-dimensional configuration sum in Sage:

```
sage: K = crystals.KirillovReshetikhin(['A',2,1],1,1)
sage: T = crystals.TensorProduct(K,K)
sage: T.one_dimensional_configuration_sum()
B[-2*Lambda[1] + 2*Lambda[2]] + (q+1)*B[-Lambda[1]]
+ (q+1)*B[Lambda[1] - Lambda[2]] + B[2*Lambda[1]]
+ B[-2*Lambda[2]] + (q+1)*B[Lambda[2]]
```

Affine Highest Weight Crystals

Affine highest weight crystals are infinite-dimensional. Their underlying weight lattice is the extended weight lattice including the null root δ . This is in contrast to finite-dimensional affine crystals such as for example the Kirillov-Reshetikhin crystals whose underlying weight lattice is the classical weight lattice, i.e., does not include δ .

Hence, to work with them in Sage, we need some further tools.

Littelmann path model

The Littelmann path model for highest weight crystals is implemented in Sage. It models finite highest crystals as well as affine highest weight crystals which are infinite dimensional. The elements of the crystal are piecewise linear maps in the extended weight space over **Q**. For more information on the Littelmann path model, see [L1995].

Since the affine highest weight crystals are infinite, it is not possible to list all elements or draw the entire crystal graph. However, if the user is only interested in the crystal up to a certain distance or depth from the highest weight element, then one can work with the corresponding subcrystal. To view the corresponding upper part of the crystal, one can build the associated digraph:

```
sage: R = RootSystem(['C', 3, 1])
sage: La = R.weight_space(extended = True).basis()
sage: LS = crystals.LSPaths(2*La[1]); LS
The crystal of LS paths of type ['C', 3, 1] and weight 2*Lambda[1]
sage: LS.weight_lattice_realization()
Extended weight space over the Rational Field of the Root system of type ['C', 3, 1]
sage: C = LS.subcrystal(max_depth=3)
sage: sorted(C, key=str)
[(-Lambda[0] + 2*Lambda[1] - Lambda[2] + Lambda[3] - delta, Lambda[1]),
 (-Lambda[0] + Lambda[1] + Lambda[2] - delta, Lambda[0] - Lambda[1] + Lambda[2]),
 (-Lambda[0] + Lambda[1] + Lambda[2] - delta, Lambda[1]),
 (-Lambda[1] + 2*Lambda[2] - delta, Lambda[1]),
 (2*Lambda[0] - 2*Lambda[1] + 2*Lambda[2],),
 (2*Lambda[1],),
 (Lambda[0] + Lambda[2] - Lambda[3], Lambda[1]),
 (Lambda[0] - Lambda[1] + Lambda[2], Lambda[1]),
 (Lambda[0] - Lambda[2] + Lambda[3], Lambda[0] - Lambda[1] + Lambda[2]),
 (Lambda[0] - Lambda[2] + Lambda[3], Lambda[1])]
sage: G = LS.digraph(subset = C)
sage: view(G, tightpage=True) # optional - dot2tex graphviz, not tested (opens external window)
                            [-\Lambda_0 + \Lambda_1 + \Lambda_2 - \delta, \Lambda_0 - \Lambda_1 + \Lambda_2] \quad [\Lambda_0 - \Lambda_2 + \Lambda_3, \Lambda_0 - \Lambda_1 + \Lambda_2] \quad [\Lambda_0 + \Lambda_2 - \Lambda_3, \Lambda_1] \quad [-\Lambda_0 + 2\Lambda_1 - \Lambda_2 + \Lambda_3 - \delta, \Lambda_1] \quad [-\Lambda_1 + 2\Lambda_2 - \delta, \Lambda_1]
```

The Littelmann path model also lends itself as a model for level zero crystals which are bi-infinite. To cut out a slice of these crystals, one can use the direction option in subcrystal:

```
sage: R = RootSystem(['A',2,1])
sage: La = R.weight_space(extended = True).basis()
sage: LS = crystals.LSPaths(La[1]-La[0]); LS
The crystal of LS paths of type ['A', 2, 1] and weight -Lambda[0] + Lambda[1]
sage: C = LS.subcrystal(max_depth=2, direction = 'both')
sage: G = LS.digraph(subset = C)
sage: G.set_latex_options(edge_options = lambda (u,v,label): ({}))
sage: view(G, tightpage=True) # optional - dot2tex graphviz, not tested (opens external window)
```



Modified Nakajima monomials

Modified Nakajima monomials have also been implemented in Sage. They model highest weight crystals in all symmetrizable types. The elements are given in terms of commuting variables $Y_i(n)$ where $i \in I$ and $n \in \mathbb{Z}_{\geq 0}$. For more information on the modified Nakajima monomials, see [KKS2007].

We give an example in affine type and verify that up to depth 3, it agrees with the Littelmann path model. Unlike in the LS path model, the Nakajima monomial crystals are indexed by elements in the weight lattice (rather than the weight space):

```
sage: La = RootSystem(['C',3,1]).weight_space(extended = True).fundamental_weights()
sage: LS = crystals.LSPaths(2*La[1]+La[2])
sage: SL = LS.subcrystal(max_depth=3)
sage: GL = LS.digraph(subset=SL)

sage: La = RootSystem(['C',3,1]).weight_lattice(extended = True).fundamental_weights()
sage: M = crystals.NakajimaMonomials(['C',3,1], 2*La[1]+La[2])
sage: SM = M.subcrystal(max_depth=3)
sage: GM = M.digraph(subset=SM)
sage: GL.is_isomorphic(GM, edge_labels=True)
True
```

Now we do an example of a simply-laced (and hence symmetrizable) hyperbolic type $H_1^{(4)}$, which comes from the complete graph on 4 vertices:

```
sage: CM = CartanMatrix([[2, -1, -1,-1],[-1,2,-1,-1],[-1,-1,2,-1],[-1,-1,-1,2]]); CM
[ 2 -1 -1 -1]
[-1  2 -1 -1]
[-1 -1  2 -1]
[-1 -1 -1  2]
sage: La = RootSystem(CM).weight_lattice().fundamental_weights()
sage: M = crystals.NakajimaMonomials(CM, La[0])
sage: SM = M.subcrystal(max_depth=4)
sage: GM = M.digraph(subset=SM) # long time
```

Elementary crystals

Note: Each of these crystals will work with any Cartan matrix input (with weights from the weight lattice corre-

sponding to the Cartan matrix given).

T-crystal

Let λ be a weight. As defined in [Kashiwara1993] (see, also, [Kashiwara1995]) the crystal $T_{\lambda} = \{t_{\lambda}\}$ is a single element crystal with the crystal structure defined by

$$\operatorname{wt}(t_{\lambda}) = \lambda, \quad e_i t_{\lambda} = f_i t_{\lambda} = 0, \quad \varepsilon_i(t_{\lambda}) = \varphi_i(t_{\lambda}) = -\infty.$$

The crystal T_{λ} shifts the weights of the vertices in a crystal B by λ when tensored with B, but leaves the graph structure of B unchanged. That is, for all $b \in B$, we have $\operatorname{wt}(t_{\lambda} \otimes b) = \operatorname{wt}(b) + \lambda$:

```
sage: B = crystals.Tableaux(['A',2],shape=[2,1])
sage: T = crystals.elementary.T(['A',2], B.Lambda()[1] + B.Lambda()[2])
sage: V = crystals.TensorProduct(T,B)
sage: for x in V:
         print(x.weight())
. . . . :
. . . . :
(4, 2, 0)
(3, 3, 0)
(3, 2, 1)
(3, 1, 2)
(2, 2, 2)
(4, 1, 1)
(3, 2, 1)
(2, 3, 1)
sage: for x in B:
. . . . :
          print(x.weight() + T[0].weight())
. . . . :
(4, 2, 0)
(3, 3, 0)
(3, 2, 1)
(3, 1, 2)
(2, 2, 2)
(4, 1, 1)
(3, 2, 1)
(2, 3, 1)
```

Warning: Sage uses the opposite convention for the tensor product rule to Kashiwara's definition, so care must be taken when comparing the examples here with Kashiwara's papers.

Here is an example using a hyperbolic Cartan matrix:

```
sage: A = CartanMatrix([[2,-4],[-4,2]])
sage: La = RootSystem(A).weight_lattice().fundamental_weights()
sage: La
Finite family {0: Lambda[0], 1: Lambda[1]}
sage: T = crystals.elementary.T(A, La[1])
sage: T
The T crystal of type [ 2 -4]
[-4 2] and weight Lambda[1]
```

C-crystal

Defined in [Kashiwara1993], the component crystal $C = \{c\}$ is the single element crystal whose crystal structure is defined by

$$\operatorname{wt}(c) = 0$$
, $e_i c = f_i c = 0$, $\varepsilon_i(c) = \varphi_i(c) = 0$.

Note $C \cong B(0)$, where B(0) is the highest weight crystal of highest weight 0.

The crystal $C \otimes T_{\mu}$ is useful when finding subcrystals inside irreducible highest weight crystals $B(\lambda)$ where λ is larger than μ in the lexicographic order. For example:

```
sage: La = RootSystem("C2").weight_lattice().fundamental_weights()
sage: h = RootSystem("C2").weight_lattice().simple_coroots()
sage: T = crystals.elementary.T("C2",2*La[1])
sage: C = crystals.elementary.Component("C2")
sage: B = crystals.TensorProduct(C,T)
sage: b = B(C[0],T[0])
sage: for i in B.index_set(): print(b.epsilon(i))
0
sage: for i in B.index_set(): print(b.phi(i))
0
sage: for i in B.index_set(): print(b.f(i))
None
None
sage: for i in B.index_set(): print(b.e(i))
None
None
```

This new crystal can be summarized into the R-crystal below.

R-crystal

For a fixed weight λ , the crystal $R_{\lambda} = \{r_{\lambda}\}$ is a single element crystal with the crystal structure defined by

$$\operatorname{wt}(r_{\lambda}) = \lambda, \quad e_{i}r_{\lambda} = f_{i}r_{\lambda} = 0, \quad \varepsilon_{i}(r_{\lambda}) = -\langle h_{i}, \lambda \rangle, \quad \varphi_{i}(r_{\lambda}) = 0,$$

where $\{h_i\}$ are the simple coroots. See page 146 [Joseph1995], for example, for more details. (Note that in [Joseph1995], this crystal is denoted by S_{λ} .)

Tensoring R_{λ} with a crystal B results in shifting the weights of the vertices in B by λ and may also cut a subset out of the original graph of B.

Warning: Sage uses the opposite convention for the tensor product rule to Kashiwara's definition, so care must be taken when comparing the examples here with some of the literature.

For example, suppose $\mu \leq \lambda$ in lexicographic ordering on weights, and one wants to see $B(\mu)$ as a subcrystal of $B(\lambda)$. Then $B(\mu)$ may be realized as the connected component of $R_{\mu-\lambda}\otimes B(\lambda)$ containing the highest weight $r_{\mu-\lambda}\otimes u_{\lambda}$, where u_{λ} is the highest weight vector in $B(\lambda)$:

```
sage: La = RootSystem(['B',4]).weight_lattice().fundamental_weights()
sage: Bla = crystals.NakajimaMonomials(['B',4], La[1]+La[2])
sage: Bmu = crystals.NakajimaMonomials(['B',4], La[1])
sage: R = crystals.elementary.R(['B',4], -La[2])
sage: T = crystals.TensorProduct(R,Bla)
```

```
sage: mg = mg = T(R[0], Bla.module_generators[0])
sage: S = T.subcrystal(generators=[mg])
sage: G = T.digraph(subset=S)
sage: Bmu.digraph().is_isomorphic(G, edge_labels=True)
True
sage: view(G, tightpage=True) # optional - dot2tex graphviz, not tested (opens external window)
```



i-th elementary crystal

For i an element of the index set of type X, the crystal B_i of type X is the set

$$B_i = \{b_i(m) : m \in \mathbf{Z}\},\$$

where the crystal stucture is given by $\operatorname{wt}(b_i(m)) = m\alpha_i$ and

$$\varphi_{j}\big(b_{i}(m)\big) = \begin{cases} m & \text{if } j = i, \\ -\infty & \text{if } j \neq i, \end{cases} \qquad \varepsilon_{j}\big(b_{i}(m)\big) = \begin{cases} -m & \text{if } j = i, \\ -\infty & \text{if } j \neq i, \end{cases}$$
$$f_{j}b_{i}(m) = \begin{cases} b_{i}(m-1) & \text{if } j = i, \\ 0 & \text{if } j \neq i, \end{cases} \qquad e_{j}b_{i}(m) = \begin{cases} b_{i}(m+1) & \text{if } j = i, \\ 0 & \text{if } j \neq i. \end{cases}$$

See [Kashiwara1993] or [Kashiwara1995] for more information. Here is an example:

```
sage: B = crystals.elementary.Elementary("A2",1)
sage: S = B.subcrystal(max_depth=4, generators=[B(0)])
sage: [s for s in S]
[0, 1, -1, 2, -2, 3, -3, -4, 4]
```

```
sage: G = B.digraph(subset=S)
sage: view(G, tightpage=True) # optional - dot2tex graphviz, not tested (opens external window)
```



Warning: To reiterate, Sage uses the opposite convention for the tensor product rule to Kashiwara's definition. In particular, using Sage's convention, one has $T_{\lambda} \otimes B_i \cong B_i \otimes T_{s_i\lambda}$, where s_i is the *i*-th simple reflection.

Infinity Crystals

Infinity crystals are the crystal analogue of Verma modules with highest weight 0 associated to a symmetrizable Kac-Moody algebra. As such, they are infinite-dimensional and any irreducible highest weight crystal may be obtained from an infinity crystal via some cutting procedure. On the other hand, the crystal $B(\infty)$ is the direct limit of all irreducible highest weight crystals $B(\lambda)$, so there are natural embeddings of each $B(\lambda)$ in $B(\infty)$. Below, we outline the various implementations of the crystal $B(\infty)$ in Sage and give examples of how $B(\lambda)$ interacts with $B(\infty)$.

All infinity crystals that are currently implemented in Sage can be accessed by typing crystals.infinity.<tab>.

Marginally large tableaux

Marginally large tableaux were introduced by J. Hong and H. Lee as a realization of the crystal $B(\infty)$ in types A_n , B_n , C_n , D_{n+1} , and G_2 . The marginally large condition guarantees that all tableau have exactly n rows and that the number of i-boxes in the i-th row from the top (in the English convention) is exactly one more than the total number of boxes in the (i+1)-st row. Other specific conditions on the tableaux vary by type. See [HongLee2008] for more information.

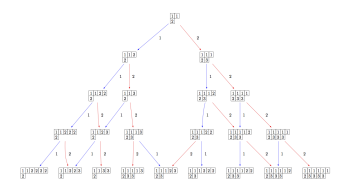
Here is an example in type A_2 :

```
sage: B = crystals.infinity.Tableaux(['A',2])
sage: b = B.highest_weight_vector()
sage: b.pp()

1  1
2
sage: b.f_string([1,2,2,1,2,1,2,2,2,2,2]).pp()
1  1  1  1  1  1  1  1  2  2  3
2  3  3  3  3  3  3  3
```

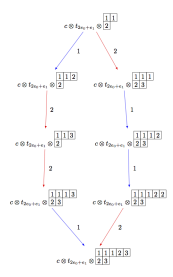
Since the crystal is infinite, we must specify the subcrystal we would like to view:

```
sage: B = crystals.infinity.Tableaux(['A',2])
sage: S = B.subcrystal(max_depth=4)
sage: G = B.digraph(subset=S)
sage: view(G, tightpage=True) # optional - dot2tex graphviz, not tested (opens external window)
```



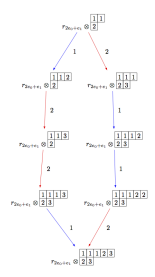
Using elementary crystals, we can cut irreducible highest weight crystals from $B(\infty)$ as the connected component of $C \otimes T_{\lambda} \otimes B(\infty)$ containing $c \otimes t_{\lambda} \otimes u_{\infty}$, where u_{∞} is the highest weight vector in $B(\infty)$. In this example, we cut out $B(\rho)$ from $B(\infty)$ in type A_2 :

```
sage: B = crystals.infinity.Tableaux(['A',2])
sage: b = B.highest_weight_vector()
sage: T = crystals.elementary.T(['A',2], B.Lambda()[1] + B.Lambda()[2])
sage: t = T[0]
sage: C = crystals.elementary.Component(['A',2])
sage: c = C[0]
sage: TP = crystals.TensorProduct(C,T,B)
sage: t0 = TP(c,t,b)
sage: STP = TP.subcrystal(generators=[t0])
sage: GTP = TP.digraph(subset=STP)
sage: view(GTP, tightpage=True) # optional - dot2tex graphviz, not tested (opens external window)
```



Note that the above code can be simplified using the R-crystal:

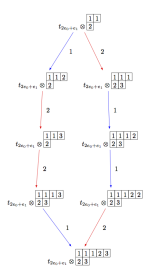
```
sage: B = crystals.infinity.Tableaux(['A',2])
sage: b = B.highest_weight_vector()
sage: R = crystals.elementary.R(['A',2], B.Lambda()[1] + B.Lambda()[2])
sage: r = R[0]
sage: TP2 = crystals.TensorProduct(R,B)
sage: t2 = TP2(r,b)
sage: STP2 = TP2.subcrystal(generators=[t2])
sage: GTP2 = TP2.digraph(subset=STP2)
sage: view(GTP2, tightpage=True) # optional - dot2tex graphviz, not tested (opens external window)
```



On the other hand, we can embed the irreducible highest weight crystal $B(\rho)$ into $B(\infty)$:

```
sage: Brho = crystals.Tableaux(['A',2],shape=[2,1])
sage: brho = Brho.highest_weight_vector()
sage: B = crystals.infinity.Tableaux(['A',2])
sage: binf = B.highest_weight_vector()
sage: wt = brho.weight()
sage: T = crystals.elementary.T(['A',2],wt)
sage: TlambdaBinf = crystals.TensorProduct(T,B)
sage: hw = TlambdaBinf(T[0],binf)
```

```
sage: Psi = Brho.crystal_morphism({brho : hw})
sage: BG = B.digraph(subset=[Psi(x) for x in Brho])
sage: view(BG, tightpage=True) # optional - dot2tex graphviz, not tested (opens external window)
```



Note that in the last example, we had to inject $B(\rho)$ into the tensor product $T_{\lambda} \otimes B(\infty)$, since otherwise, the map Psi would not be a crystal morphism (as b.weight () != brho.weight ()).

Generalized Young walls

Generalized Young walls were introduced by J.-A. Kim and D.-U. Shin as a model for $B(\infty)$ and each $B(\lambda)$ solely in affine type $A_n^{(1)}$. See [KimShin2010] for more information on the construction of generalized Young walls.

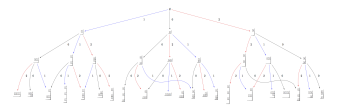
Since this model is only valid for one Cartan type, the input to initialize the crystal is simply the rank of the underlying type:

In the weight method for this model, we can choose whether to view weights in the extended weight lattice (by default) or in the root lattice:

```
sage: Y = crystals.infinity.GeneralizedYoungWalls(2)
sage: y = Y.highest_weight_vector()
sage: y.f_string([0,1,2,2,2,1,0,0,1,2]).weight()
Lambda[0] + Lambda[1] - 2*Lambda[2] - 3*delta
sage: y.f_string([0,1,2,2,2,1,0,0,1,2]).weight(root_lattice=True)
-3*alpha[0] - 3*alpha[1] - 4*alpha[2]
```

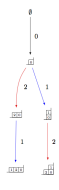
As before, we need to indicate a specific subcrystal when attempting to view the crystal graph:

```
sage: Y = crystals.infinity.GeneralizedYoungWalls(2)
sage: SY = Y.subcrystal(max_depth=3)
sage: GY = Y.digraph(subset=SY)
sage: view(GY, tightpage=True) # optional - dot2tex graphviz, not tested (opens external window)
```



One can also make irreducible highest weight crystals using generalized Young walls:

```
sage: La = RootSystem(['A',2,1]).weight_lattice(extended=True).fundamental_weights()
sage: YLa = crystals.GeneralizedYoungWalls(2,La[0])
sage: SYLa = YLa.subcrystal(max_depth=3)
sage: GYLa = YLa.digraph(subset=SYLa)
sage: view(GYLa, tightpage=True) # optional - dot2tex graphviz, not tested (opens external window)
```



In the highest weight crystals, however, weights are always elements of the extended affine weight lattice:

```
sage: La = RootSystem(['A',2,1]).weight_lattice(extended=True).fundamental_weights()
sage: YLa = crystals.GeneralizedYoungWalls(2,La[0])
sage: YLa.highest_weight_vector().f_string([0,1,2,0]).weight()
-Lambda[0] + Lambda[1] + Lambda[2] - 2*delta
```

Modified Nakajima monomials

Let $Y_{i,k}$, for $i \in I$ and $k \in \mathbf{Z}$, be a commuting set of variables, and let 1 be a new variable which commutes with each $Y_{i,k}$. (Here, I represents the index set of a Cartan datum.) One may endow the structure of a crystal on the set $\widehat{\mathcal{M}}$ of monomials of the form

$$M = \prod_{(i,k) \in I \times \mathbf{Z}_{\geq 0}} Y_{i,k}^{y_i(k)} \mathbf{1}.$$

Elements of $\widehat{\mathcal{M}}$ are called *modified Nakajima monomials*. We will omit the 1 from the end of a monomial if there exists at least one $y_i(k) \neq 0$. The crystal structure on this set is defined by

$$\operatorname{wt}(M) = \sum_{i \in I} \left(\sum_{k \geq 0} y_i(k) \right) \Lambda_i,$$

$$\varphi_i(M) = \max \left\{ \sum_{0 \leq j \leq k} y_i(j) : k \geq 0 \right\},$$

$$\varepsilon_i(M) = \varphi_i(M) - \langle h_i, \operatorname{wt}(M) \rangle,$$

$$k_f = k_f(M) = \min \left\{ k \geq 0 : \varphi_i(M) = \sum_{0 \leq j \leq k} y_i(j) \right\},$$

$$k_e = k_e(M) = \max \left\{ k \geq 0 : \varphi_i(M) = \sum_{0 \leq j \leq k} y_i(j) \right\},$$

where $\{h_i : i \in I\}$ and $\{\Lambda_i : i \in I\}$ are the simple coroots and fundamental weights, respectively. With a chosen set of integers $C = (c_{ij})_{i \neq j}$ such that $c_{ij} + cji = 1$, one defines

$$A_{i,k} = Y_{i,k} Y_{i,k+1} \prod_{j \neq i} Y_{j,k+c_{ji}}^{a_{ji}},$$

where (a_{ij}) is a Cartan matrix. Then

$$e_i M = \begin{cases} 0 & \text{if } \varepsilon_i(M) = 0, \\ A_{i,k_e} M & \text{if } \varepsilon_i(M) > 0, \end{cases}$$
$$f_i M = A_{i,k_f}^{-1} M.$$

Note: Monomial crystals depend on the choice of positive integers $C = (c_{ij})_{i \neq j}$ satisfying the condition $c_{ij} + c_{ji} = 1$. This choice has been made in Sage such that $c_{ij} = 1$ if i < j and $c_{ij} = 0$ if i > j.

It is shown in [KKS2007] that the connected component of $\widehat{\mathcal{M}}$ containing the element 1, which we denote by $\mathcal{M}(\infty)$, is crystal isomorphic to the crystal $B(\infty)$:

```
sage: Minf = crystals.infinity.NakajimaMonomials(['C',3,1])
sage: minf = Minf.highest_weight_vector()
sage: m = minf.f_string([0,1,2,3,2,1,0]); m
Y(0,0)^-1 Y(0,4)^-1 Y(1,0) Y(1,3)
sage: m.weight()
-2*Lambda[0] + 2*Lambda[1] - 2*delta
sage: m.weight_in_root_lattice()
-2*alpha[0] - 2*alpha[1] - 2*alpha[2] - alpha[3]
```

We can also model $B(\infty)$ using the monomials $A_{i,k}$ instead:

```
sage: Ninf = crystals.infinity.NakajimaMonomials(['C',3,1], use_Y = False)
sage: ninf = Ninf.highest_weight_vector()
sage: n = ninf.f_string([0,1,2,3,2,1,0]); n
A(0,0)^-1 A(0,3)^-1 A(1,0)^-1 A(1,2)^-1 A(2,0)^-1 A(2,1)^-1 A(3,0)^-1
sage: n.weight()
-2*Lambda[0] + 2*Lambda[1] - 2*delta
sage: n.weight_in_root_lattice()
-2*alpha[0] - 2*alpha[1] - 2*alpha[2] - alpha[3]
```

Building the crystal graph output for these monomial crystals is the same as the constructions above:

```
sage: Minf = crystals.infinity.NakajimaMonomials(['C',3,1])
sage: Sinf = Minf.subcrystal(max_depth=2)
sage: Ginf = Minf.digraph(subset=Sinf)
sage: view(Ginf, tightpage=True) # optional - dot2tex graphviz, not tested (opens external window)
```

TOTAL TRANSPORT TOTAL TO

Note that this model will also work for any symmetrizable Cartan matrix:

```
sage: A = CartanMatrix([[2,-4],[-5,2]])
sage: Linf = crystals.infinity.NakajimaMonomials(A); Linf
Infinity Crystal of modified Nakajima monomials of type [ 2 -4]
[-5 2]
sage: Linf.highest_weight_vector().f_string([0,1,1,1,0,0,1,1,0])
Y(0,0)^{-1} Y(0,1)^9 Y(0,2)^5 Y(0,3)^{-1} Y(1,0)^2 Y(1,1)^5 Y(1,2)^3
```

Rigged configurations

Rigged configurations are sequences of partitions, one partition for each node in the underlying Dynkin diagram, such that each part of each partition has a label (or rigging). A crystal structure was defined on these objects in [Schilling2006], then later extended to work as a model for $B(\infty)$. See [SalisburyScrimshaw2015] for more information:

We can check this crystal is isomorphic to the crystal above using Nakajima monomials:

```
sage: Minf = crystals.infinity.NakajimaMonomials(['C',3,1])
sage: Sinf = Minf.subcrystal(max_depth=2)
sage: Ginf = Minf.digraph(subset=Sinf)
sage: RC = crystals.infinity.RiggedConfigurations(['C',3,1])
sage: RCS = RC.subcrystal(max_depth=2)
sage: RCG = RC.digraph(subset=RCS)
sage: RCG.is_isomorphic(Ginf, edge_labels=True)
True
```

This model works in Sage for all finite and affine types, as well as any simply laced Cartan matrix.

Iwahori Hecke Algebras

The Iwahori Hecke algebra is defined in [Iwahori1964]. In that original paper, the algebra occurs as the convolution ring of functions on a p-adic group that are compactly supported and invariant both left and right by the Iwahori subgroup. However Iwahori determined its structure in terms of generators and relations, and it turns out to be a deformation of the group algebra of the affine Weyl group.

Once the presentation is found, the Iwahori Hecke algebra can be defined for any Coxeter group. It depends on a parameter q which in Iwahori's paper is the cardinality of the residue field. But it could just as easily be an indeterminate.

Then the Iwahori Hecke algebra has the following description. Let W be a Coxeter group, with generators (simple reflections) s_1, \ldots, s_n . They satisfy the relations $s_i^2 = 1$ and the braid relations

$$s_i s_j s_i s_j \cdots = s_j s_i s_j s_i \cdots$$

where the number of terms on each side is the order of $s_i s_j$.

The Iwahori Hecke algebra has a basis T_1, \ldots, T_n subject to relations that resemble those of the s_i . They satisfy the braid relations and the quadratic relation

$$(T_i - q)(T_i + 1) = 0.$$

This can be modified by letting q_1 and q_2 be two indeterminates and letting

$$(T_i - q_1)(T_i - q_2) = 0.$$

In this generality, Iwahori Hecke algebras have significance far beyond their origin in the representation theory of *p*-adic groups. For example, they appear in the geometry of Schubert varieties, where they are used in the definition of the Kazhdan-Lusztig polynomials. They appear in connection with quantum groups, and in Jones's original paper on the Jones polynomial.

Here is how to create an Iwahori Hecke algebra (in the T basis):

```
sage: R.<q> = PolynomialRing(ZZ)
sage: H = IwahoriHeckeAlgebra("B3",q)
sage: T = H.T(); T
Iwahori-Hecke algebra of type B3 in q,-1 over Univariate Polynomial Ring
in q over Integer Ring in the T-basis
sage: T1,T2,T3 = T.algebra_generators()
sage: T1*T1
(q-1)*T[1] + q
```

If the Cartan type is affine, the generators will be numbered starting with T0 instead of T1.

You may coerce a Weyl group element into the Iwahori Hecke algebra:

```
sage: W = WeylGroup("G2",prefix="s")
sage: [s1,s2] = W.simple_reflections()
sage: P.<q> = LaurentPolynomialRing(QQ)
sage: H = IwahoriHeckeAlgebra("B3",q)
sage: T = H.T()
sage: T(s1*s2)
T[1,2]
```

Kazhdan-Lusztig Polynomials

Sage can compute ordinary Kazhdan-Lusztig polynomials for Weyl groups or affine Weyl groups (and potentially other Coxeter groups).

You must create a Weyl group w and a ring containing an indeterminate q. The ring may be a univariate polynomial ring or a univariate Laurent polynomial ring. Then you may calculate Kazhdan-Lusztig polynomials as follows:

```
sage: W = WeylGroup("A3", prefix="s")
sage: [s1,s2,s3] = W.simple_reflections()
sage: P.<q> = LaurentPolynomialRing(QQ)
sage: KL = KazhdanLusztigPolynomial(W,q)
sage: KL.R(s2, s2*s1*s3*s2)
-1 + 3*q - 3*q^2 + q^3
sage: KL.P(s2, s2*s1*s3*s2)
1 + q
```

Thus we have the Kazhdan-Lusztig R and P polynomials.

Known algorithms for computing Kazhdan-Lusztig polynomials are highly recursive, and caching of intermediate results is necessary for the programs not to be prohibitively slow. Therefore intermediate results are cached. This has the effect that as you run the program for any given KazhdanLusztigPolynomial class, the calculations will be slow at first but progressively faster as more polynomials are computed.

You may see the results of the intermediate calculations by creating the class with the option trace="true".

Since the parent of q must be a univariate ring, if you want to work with other indeterminates, *first* create a univariate polynomial or Laurent polynomial ring, and the Kazhdan-Lusztig class. *Then* create a ring containing q and the other variables:

```
sage: W = WeylGroup("B3", prefix="s")
sage: [s1,s2,s3] = W.simple_reflections()
sage: P.<q> = PolynomialRing(QQ)
sage: KL = KazhdanLusztigPolynomial(W,q)
sage: P1.<x,y> = PolynomialRing(P)
sage: x*KL.P(s1*s3,s1*s3*s2*s1*s3)
(q + 1)*x
```

Bibliography

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12.1.6 Linear Programming (Mixed Integer)

This document explains the use of linear programming (LP) – and of mixed integer linear programming (MILP) – in Sage by illustrating it with several problems it can solve. Most of the examples given are motivated by graph-theoretic concerns, and should be understandable without any specific knowledge of this field. As a tool in Combinatorics, using linear programming amounts to understanding how to reformulate an optimization (or existence) problem through linear constraints.

This is a translation of a chapter from the book Calcul mathematique avec Sage.

Definition

Here we present the usual definition of what a linear program is: it is defined by a matrix $A : \mathbb{R}^m \to \mathbb{R}^n$, along with two vectors $b, c \in \mathbb{R}^n$. Solving a linear program is searching for a vector x maximizing an *objective* function and satisfying a set of constraints, i.e.

$$c^t x = \max_{x' \text{ such that } Ax' \le b} c^t x'$$

where the ordering $u \le u'$ between two vectors means that the entries of u' are pairwise greater than the entries of u. We also write:

$$\begin{aligned} & \text{Max: } c^t x \\ & \text{Such that: } Ax \leq b \end{aligned}$$

Equivalently, we can also say that solving a linear program amounts to maximizing a linear function defined over a polytope (preimage or $A^{-1}(\leq b)$). These definitions, however, do not tell us how to use linear programming in combinatorics. In the following, we will show how to solve optimization problems like the Knapsack problem, the Maximum Matching problem, and a Flow problem.

Mixed integer linear programming

There are bad news coming along with this definition of linear programming: an LP can be solved in polynomial time. This is indeed bad news, because this would mean that unless we define LP of exponential size, we cannot expect LP to solve NP-complete problems, which would be a disappointment. On a brighter side, it becomes NP-complete to solve a linear program if we are allowed to specify constraints of a different kind: requiring that some variables be integers instead of real values. Such an LP is actually called a "mixed integer linear program" (some variables can be integers, some other reals). Hence, we can expect to find in the MILP framework a *wide* range of expressivity.

Practical

The MILP class

The MILP class in Sage represents a MILP! It is also used to solve regular LP. It has a very small number of methods, meant to define our set of constraints and variables, then to read the solution found by the solvers once computed. It is also possible to export a MILP defined with Sage to a .lp or .mps file, understood by most solvers.

Let us ask Sage to solve the following LP:

```
Max: x + y + 3z
Such that: x + 2y \le 4
5z - y \le 8
x, y, z \ge 0
```

To achieve it, we need to define a corresponding MILP object, along with 3 variables x, y and z:

```
sage: p = MixedIntegerLinearProgram()
sage: v = p.new_variable(real=True, nonnegative=True)
sage: x, y, z = v['x'], v['y'], v['z']
```

Next, we set the objective function

```
sage: p.set_objective(x + y + 3*z)
```

And finally we set the constraints

```
sage: p.add_constraint(x + 2*y <= 4)
sage: p.add_constraint(5*z - y <= 8)</pre>
```

The solve method returns by default the optimal value reached by the objective function

```
sage: round(p.solve(), 2)
8.8
```

We can read the optimal assignation found by the solver for x, y and z through the get_values method

```
sage: round(p.get_values(x), 2)
4.0
sage: round(p.get_values(y), 2)
0.0
sage: round(p.get_values(z), 2)
1.6
```

Variables

In the previous example, we obtained variables through v['x'], v['y'] and v['z']. This being said, larger LP/MILP will require us to associate an LP variable to many Sage objects, which can be integers, strings, or even the vertices and edges of a graph. For example:

```
sage: x = p.new_variable(real=True, nonnegative=True)
```

With this new object x we can now write constraints using $x[1], \ldots, x[15]$.

```
sage: p.add_constraint(x[1] + x[12] - x[14] >= 8)
```

Notice that we did not need to define the "length" of x. Actually, x would accept any immutable object as a key, as a dictionary would. We can now write

And because any immutable object can be used as a key, doubly indexed variables $x^{1,1}, ..., x^{1,15}, x^{2,1}, ..., x^{15,15}$ can be referenced by x[1,1], ..., x[1,15], x[2,1], ..., x[15,15]

```
sage: p.add_constraint(x[3,2] + x[5] == 6)
```

Typed variables and bounds Types: If you want a variable to assume only integer or binary values, use the integer=True or binary=True arguments of the new_variable method. Alternatively, call the set integer and set binary methods.

Bounds: If you want your variables to only take nonnegative values, you can say so when calling new_variable with the argument nonnegative=True. If you want to set a different upper/lower bound on a variable, add a constraint or use the set_min, set_max methods.

Basic linear programs

Knapsack

The *Knapsack* problem is the following: given a collection of items having both a weight and a *usefulness*, we would like to fill a bag whose capacity is constrained while maximizing the usefulness of the items contained in the bag (we will consider the sum of the items' usefulness). For the purpose of this tutorial, we set the restriction that the bag can only carry a certain total weight.

To achieve this, we have to associate to each object o of our collection C a binary variable taken [o], set to 1 when the object is in the bag, and to 0 otherwise. We are trying to solve the following MILP

```
\begin{aligned} & \text{Max: } \sum_{o \in L} \text{usefulness}_o \times \text{taken}_o \\ & \text{Such that: } \sum_{o \in L} \text{weight}_o \times \text{taken}_o \leq C \end{aligned}
```

Using Sage, we will give to our items a random weight:

```
sage: C = 1
sage: L = ["pan", "book", "knife", "gourd", "flashlight"]
```

```
sage: L.extend(["random_stuff_" + str(i) for i in range(20)])
sage: weight = {}
sage: usefulness = {}
sage: set_random_seed(685474)
sage: for o in L:
        weight[o] = random()
. . . . :
          usefulness[0] = random()
We can now define the MILP itself
sage: p = MixedIntegerLinearProgram()
sage: taken = p.new_variable(binary=True)
sage: p.add_constraint(sum(weight[o] * taken[o] for o in L) <= C)</pre>
sage: p.set_objective(sum(usefulness[o] * taken[o] for o in L))
sage: p.solve() # abs tol 1e-6
3.1502766806530307
sage: taken = p.get_values(taken)
The solution found is (of course) admissible
sage: sum(weight[o] * taken[o] for o in L) # abs tol 1e-6
0.6964959796619171
Should we take a flashlight?
sage: taken["flashlight"]
1.0
```

Wise advice. Based on purely random considerations.

Matching

Given a graph G, a matching is a set of pairwise disjoint edges. The empty set is a trivial matching. So we focus our attention on maximum matchings: we want to find in a graph a matching whose cardinality is maximal. Computing the maximum matching in a graph is a polynomial problem, which is a famous result of Edmonds. Edmonds' algorithm is based on local improvements and the proof that a given matching is maximum if it cannot be improved. This algorithm is not the hardest to implement among those graph theory can offer, though this problem can be modeled with a very simple MILP.

To do it, we need – as previously – to associate a binary variable to each one of our objects: the edges of our graph (a value of 1 meaning that the corresponding edge is included in the maximum matching). Our constraint on the edges taken being that they are disjoint, it is enough to require that, x and y being two edges and m_x, m_y their associated variables, the inequality $m_x + m_y \le 1$ is satisfied, as we are sure that the two of them cannot both belong to the matching. Hence, we are able to write the MILP we want. However, the number of inequalities can be easily decreased by noticing that two edges cannot be taken simultaneously inside a matching if and only if they have a common endpoint v. We can then require instead that at most one edge incident to v be taken inside the matching,

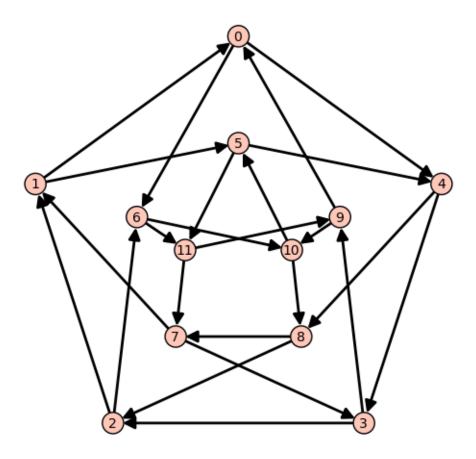
which is a linear constraint. We will be solving:

$$\begin{aligned} & \text{Max: } \sum_{e \in E(G)} m_e \\ & \text{Such that: } \forall v, \sum_{e \in E(G)} m_e \leq 1 \end{aligned}$$

Let us write the Sage code of this MILP:

Flows

Yet another fundamental algorithm in graph theory: maximum flow! It consists, given a directed graph and two vertices s, t, in sending a maximum flow from s to t using the edges of G, each of them having a maximal capacity.



The definition of this problem is almost its LP formulation. We are looking for real values associated to each edge, which would represent the intensity of flow going through them, under two types of constraints:

- The amount of flow arriving on a vertex (different from s or t) is equal to the amount of flow leaving it.
- The amount of flow going through an edge is bounded by the capacity of this edge.

This being said, we have to maximize the amount of flow leaving s: all of it will end up in t, as the other vertices are sending just as much as they receive. We can model the flow problem with the following LP

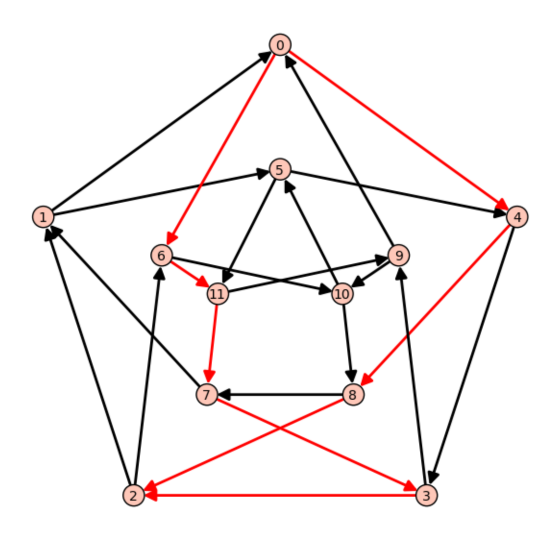
Max:
$$\sum_{sv \in G} f_{sv}$$

Such that: $\forall v \in G, \frac{v \neq s}{v \neq t}, \sum_{vu \in G} f_{vu} - \sum_{uv \in G} f_{uv} = 0$
 $\forall uv \in G, f_{uv} \leq 1$

We will solve the flow problem on an orientation of Chvatal's graph, in which all the edges have a capacity of 1:

```
sage: g = graphs.ChvatalGraph()
sage: g = g.minimum_outdegree_orientation()

sage: p = MixedIntegerLinearProgram()
sage: f = p.new_variable(real=True, nonnegative=True)
sage: s, t = 0, 2
```



Solvers

Sage solves linear programs by calling specific libraries. The following libraries are currently supported:

• CBC: A solver from COIN-OR

Provided under the open source license CPL, but incompatible with GPL. CBC can be installed using the shell command sage -i cbc sagelib.

CPLEX: A solver from ILOG

Proprietary, but free for researchers and students.

 CVXOPT: an LP solver from Python Software for Convex Optimization, uses an interior-point method, always installed in Sage.

Licensed under the GPL.

• GLPK: A solver from GNU

Licensed under the GPLv3. This solver is always installed, as the default one, in Sage.

GUROBI

Proprietary, but free for researchers and students.

• PPL: A solver from bugSeng.

This solver provides exact (arbitrary precision) computation, always installed in Sage.

Licensed under the GPLv3.

Using CPLEX or GUROBI through Sage

ILOG's CPLEX and GUROBI being proprietary softwares, you must be in possession of several files to use it through Sage. In each case, the **expected** (it may change!) filename is joined.

- · A valid license file
 - CPLEX: a .ilm file
 - GUROBI: a .lic file
- · A compiled version of the library
 - CPLEX: libcplex.a
 - GUROBI: libgurobi55.so (or more recent)
- The library file
 - CPLEX:cplex.h
 - GUROBI: gurobi_c.h

The environment variable defining the licence's path must also be set when running Sage. You can append to your .bashrc file one of the following:

For CPLEX

```
export ILOG_LICENSE_FILE=/path/to/the/license/ilog/ilm/access_1.ilm
```

· For GUROBI

```
export GRB_LICENSE_FILE=/path/to/the/license/gurobi.lic
```

As Sage also needs the files library and header files the easiest way is to create symbolic links to these files in the appropriate directories:

- For CPLEX:
 - libcplex.a-in SAGE_ROOT/local/lib/, type:

```
    libgurobi56.so - in SAGE_ROOT/local/lib/, type:
    ln -s /path/to/lib/libgurobi56.so libgurobi.so
    gurobi_c.h - in SAGE_ROOT/local/include/, type:
    ln -s /path/to/include/gurobi_c.h .
```

It is very important that the names of the symbolic links in Sage's folders ** be precisely as indicated. If the names differ, Sage will not notice that ** the files are present

Once this is done, Sage is to be asked to notice the changes by running:

make

12.1.7 Number Theory and the RSA Public Key Cryptosystem

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This tutorial uses Sage to study elementary number theory and the RSA public key cryptosystem. A number of Sage commands will be presented that help us to perform basic number theoretic operations such as greatest common divisor and Euler's phi function. We then present the RSA cryptosystem and use Sage's built-in commands to encrypt and decrypt data via the RSA algorithm. Note that this tutorial on RSA is for pedagogy purposes only. For further details on cryptography or the security of various cryptosystems, consult specialized texts such as [MenezesEtAl1996], [Stinson2006], and [TrappeWashington2006].

Elementary number theory

We first review basic concepts from elementary number theory, including the notion of primes, greatest common divisors, congruences and Euler's phi function. The number theoretic concepts and Sage commands introduced will be referred to in later sections when we present the RSA algorithm.

Prime numbers

Public key cryptography uses many fundamental concepts from number theory, such as prime numbers and greatest common divisors. A positive integer n > 1 is said to be *prime* if its factors are exclusively 1 and itself. In Sage, we can obtain the first 20 prime numbers using the command primes_first_n:

```
sage: primes_first_n(20)
[2, 3, 5, 7, 11, 13, 17, 19, 23, 29, 31, 37, 41, 43, 47, 53, 59, 61, 67, 71]
```

Greatest common divisors

Let a and b be integers, not both zero. Then the greatest common divisor (GCD) of a and b is the largest positive integer which is a factor of both a and b. We use gcd(a,b) to denote this largest positive factor. One can extend this definition by setting gcd(0,0) = 0. Sage uses gcd(a,b) to denote the GCD of a and b. The GCD of any two distinct primes is 1, and the GCD of 18 and 27 is 9.

```
sage: gcd(3, 59)
1
sage: gcd(18, 27)
9
```

If gcd(a, b) = 1, we say that a is *coprime* (or relatively prime) to b. In particular, gcd(3, 59) = 1 so 3 is coprime to 59 and vice versa.

Congruences

When one integer is divided by a non-zero integer, we usually get a remainder. For example, upon dividing 23 by 5, we get a remainder of 3; when 8 is divided by 5, the remainder is again 3. The notion of congruence helps us to describe the situation in which two integers have the same remainder upon division by a non-zero integer. Let $a, b, n \in \mathbf{Z}$ such that $n \neq 0$. If a and b have the same remainder upon division by n, then we say that a is *congruent* to b modulo n and denote this relationship by

```
a \equiv b \pmod{n}
```

This definition is equivalent to saying that n divides the difference of a and b, i.e. $n \mid (a-b)$. Thus $23 \equiv 8 \pmod{5}$ because when both 23 and 8 are divided by 5, we end up with a remainder of 3. The command mod allows us to compute such a remainder:

```
sage: mod(23, 5)
3
sage: mod(8, 5)
3
```

Euler's phi function

Consider all the integers from 1 to 20, inclusive. List all those integers that are coprime to 20. In other words, we want to find those integers n, where $1 \le n \le 20$, such that $\gcd(n, 20) = 1$. The latter task can be easily accomplished with a little bit of Sage programming:

```
sage: for n in range(1, 21):
....: if gcd(n, 20) == 1:
....: print n,
...
1 3 7 9 11 13 17 19
```

The above programming statements can be saved to a text file called, say, /home/mvngu/totient.sage, organizing it as follows to enhance readability.

```
for n in xrange(1, 21):
    if gcd(n, 20) == 1:
        print n,
```

We refer to totient.sage as a Sage script, just as one would refer to a file containing Python code as a Python script. We use 4 space indentations, which is a coding convention in Sage as well as Python programming, instead of tabs.

The command load can be used to read the file containing our programming statements into Sage and, upon loading the content of the file, have Sage execute those statements:

```
load("/home/mvngu/totient.sage")
1 3 7 9 11 13 17 19
```

From the latter list, there are 8 integers in the closed interval [1, 20] that are coprime to 20. Without explicitly generating the list

```
1 3 7 9 11 13 17 19
```

how can we compute the number of integers in [1,20] that are coprime to 20? This is where Euler's phi function comes in handy. Let $n \in \mathbf{Z}$ be positive. Then *Euler's phi function* counts the number of integers a, with $1 \le a \le n$, such that $\gcd(a,n)=1$. This number is denoted by $\varphi(n)$. Euler's phi function is sometimes referred to as Euler's totient function, hence the name totient.sage for the above Sage script. The command euler_phi implements Euler's phi function. To compute $\varphi(20)$ without explicitly generating the above list, we proceed as follows:

```
sage: euler_phi(20)
```

How to keep a secret?

Cryptography is the science (some might say art) of concealing data. Imagine that we are composing a confidential email to someone. Having written the email, we can send it in one of two ways. The first, and usually convenient, way is to simply press the send button and not care about how our email will be delivered. Sending an email in this manner is similar to writing our confidential message on a postcard and post it without enclosing our postcard inside an envelope. Anyone who can access our postcard can see our message. On the other hand, before sending our email, we can scramble the confidential message and then press the send button. Scrambling our message is similar to enclosing our postcard inside an envelope. While not 100% secure, at least we know that anyone wanting to read our postcard has to open the envelope.

In cryptography parlance, our message is called *plaintext*. The process of scrambling our message is referred to as *encryption*. After encrypting our message, the scrambled version is called *ciphertext*. From the ciphertext, we can recover our original unscrambled message via *decryption*. The following figure illustrates the processes of encryption and decryption. A *cryptosystem* is comprised of a pair of related encryption and decryption processes.

```
+ -----+ encrypt +------+ decrypt +------+ | plaintext | ---------> | ciphertext | -------> | plaintext |
```

The following table provides a very simple method of scrambling a message written in English and using only upper case letters, excluding punctuation characters.

Formally, let

$$\Sigma = \{\mathtt{A},\mathtt{B},\mathtt{C},\ldots,\mathtt{Z}\}$$

be the set of capital letters of the English alphabet. Furthermore, let

$$\Phi = \{65, 66, 67, \dots, 90\}$$

be the American Standard Code for Information Interchange (ASCII) encodings of the upper case English letters. Then the above table explicitly describes the mapping $f:\Sigma\longrightarrow\Phi$. (For those familiar with ASCII, f is actually a common process for *encoding* elements of Σ , rather than a cryptographic "scrambling" process *per se*.) To scramble a message written using the alphabet Σ , we simply replace each capital letter of the message with its corresponding ASCII encoding. However, the scrambling process described in the above table provides, cryptographically speaking, very little to no security at all and we strongly discourage its use in practice.

Keeping a secret with two keys

The Rivest, Shamir, Adleman (RSA) cryptosystem is an example of a *public key cryptosystem*. RSA uses a *public key* to encrypt messages and decryption is performed using a corresponding *private key*. We can distribute our public keys, but for security reasons we should keep our private keys to ourselves. The encryption and decryption processes draw upon techniques from elementary number theory. The algorithm below is adapted from page 165 of [TrappeWashington2006]. It outlines the RSA procedure for encryption and decryption.

- 1. Choose two primes p and q and let n = pq.
- 2. Let $e \in \mathbf{Z}$ be positive such that $\gcd(e, \varphi(n)) = 1$.
- 3. Compute a value for $d \in \mathbf{Z}$ such that $de \equiv 1 \pmod{\varphi(n)}$.
- 4. Our public key is the pair (n, e) and our private key is the triple (p, q, d).
- 5. For any non-zero integer m < n, encrypt m using $c \equiv m^e \pmod{n}$.
- 6. Decrypt c using $m \equiv c^d \pmod{n}$.

The next two sections will step through the RSA algorithm, using Sage to generate public and private keys, and perform encryption and decryption based on those keys.

Generating public and private keys

Positive integers of the form $M_m=2^m-1$ are called *Mersenne numbers*. If p is prime and $M_p=2^p-1$ is also prime, then M_p is called a *Mersenne prime*. For example, 31 is prime and $M_{31}=2^{31}-1$ is a Mersenne prime, as can be verified using the command <code>is_prime(p)</code>. This command returns <code>True</code> if its argument <code>p</code> is precisely a prime number; otherwise it returns <code>False</code>. By definition, a prime must be a positive integer, hence <code>is_prime(-2)</code> returns <code>False</code> although we know that 2 is prime. Indeed, the number $M_{61}=2^{61}-1$ is also a Mersenne prime. We can use M_{31} and M_{61} to work through step 1 in the RSA algorithm:

```
sage: p = (2^31) - 1
sage: is_prime(p)
True
sage: q = (2^61) - 1
sage: is_prime(q)
True
sage: n = p * q; n
4951760154835678088235319297
```

A word of warning is in order here. In the above code example, the choice of p and q as Mersenne primes, and with so many digits far apart from each other, is a very bad choice in terms of cryptographic security. However, we shall use the above chosen numeric values for p and q for the remainder of this tutorial, always bearing in mind that they have been chosen for pedagogy purposes only. Refer to [MenezesEtAl1996], [Stinson2006], and [TrappeWashington2006] for in-depth discussions on the security of RSA, or consult other specialized texts.

For step 2, we need to find a positive integer that is coprime to $\varphi(n)$. The set of integers is implemented within the Sage module sage.rings.integer_ring. Various operations on integers can be accessed via the ZZ.* family of functions. For instance, the command ZZ.random_element(n) returns a pseudo-random integer uniformly distributed within the closed interval [0, n-1].

We can compute the value $\varphi(n)$ by calling the sage function <code>euler_phi(n)</code>, but for arbitrarily large prime numbers p and q, this can take an enormous amount of time. Indeed, the private key can be quickly deduced from the public key once you know $\varphi(n)$, so it is an important part of the security of the RSA cryptosystem that $\varphi(n)$ cannot be computed in a short time, if only n is known. On the other hand, if the private key is available, we can compute $\varphi(n) = (p-1)(q-1)$ in a very short time.

Using a simple programming loop, we can compute the required value of e as follows:

```
sage: p = (2^31) - 1
sage: q = (2^61) - 1
sage: n = p * q
sage: phi = (p - 1)*(q - 1); phi
4951760152529835076874141700
sage: e = ZZ.random_element(phi)
sage: while gcd(e, phi) != 1:
....: e = ZZ.random_element(phi)
...
sage: e # random
1850567623300615966303954877
sage: e < n
True</pre>
```

As e is a pseudo-random integer, its numeric value changes after each execution of e = ZZ.random_element(phi).

To calculate a value for d in step 3 of the RSA algorithm, we use the extended Euclidean algorithm. By definition of congruence, $de \equiv 1 \pmod{\varphi(n)}$ is equivalent to

$$de - k \cdot \varphi(n) = 1$$

where $k \in \mathbf{Z}$. From steps 1 and 2, we already know the numeric values of e and $\varphi(n)$. The extended Euclidean algorithm allows us to compute d and -k. In Sage, this can be accomplished via the command \gcd . Given two integers x and y, $\gcd(x, y)$ returns a 3-tuple (g, s, t) that satisfies the Bézout identity $g = \gcd(x, y) = sx + ty$. Having computed a value for d, we then use the command $\gcd(d * e, phi)$ to check that d * e is indeed congruent to 1 modulo phi.

```
sage: n = 4951760154835678088235319297
sage: e = 1850567623300615966303954877
sage: phi = 4951760152529835076874141700
sage: bezout = xgcd(e, phi); bezout # random
(1, 4460824882019967172592779313, -1667095708515377925087033035)
sage: d = Integer(mod(bezout[1], phi)); d # random
4460824882019967172592779313
sage: mod(d * e, phi)
1
```

Thus, our RSA public key is

```
(n, e) = (4951760154835678088235319297, 1850567623300615966303954877)
```

and our corresponding private key is

```
(p,q,d) = (2147483647, 2305843009213693951, 4460824882019967172592779313)
```

Encryption and decryption

Suppose we want to scramble the message HELLOWORLD using RSA encryption. From the above ASCII table, our message maps to integers of the ASCII encodings as given below.

```
| H E L L O W O R L D |
| 72 69 76 76 79 87 79 82 76 68 |
```

Concatenating all the integers in the last table, our message can be represented by the integer

```
m = 72697676798779827668
```

There are other more cryptographically secure means for representing our message as an integer. The above process is used for demonstration purposes only and we strongly discourage its use in practice. In Sage, we can obtain an integer representation of our message as follows:

```
sage: m = "HELLOWORLD"
sage: m = map(ord, m); m
[72, 69, 76, 76, 79, 87, 79, 82, 76, 68]
sage: m = ZZ(list(reversed(m)), 100); m
72697676798779827668
```

To encrypt our message, we raise m to the power of e and reduce the result modulo n. The command $mod(a^b, n)$ first computes a^b and then reduces the result modulo n. If the exponent b is a "large" integer, say with more than 20 digits, then performing modular exponentiation in this naive manner takes quite some time. Brute force (or naive) modular exponentiation is inefficient and, when performed using a computer, can quickly consume a huge quantity of the computer's memory or result in overflow messages. For instance, if we perform naive modular exponentiation using the command $mod(m^e, n)$, where m, n and e are as given above, we would get an error message similar to the following:

```
mod(m^e, n)
Traceback (most recent call last)
/home/mvngu/<ipython console> in <module>()
/home/mvngu/usr/bin/sage-3.1.4/local/lib/python2.5/site-packages/sage/rings/integer.so
in sage.rings.integer.Integer.__pow__ (sage/rings/integer.c:9650)()
RuntimeError: exponent must be at most 2147483647
```

There is a trick to efficiently perform modular exponentiation, called the method of repeated squaring, cf. page 879 of [CormenEtAl2001]. Suppose we want to compute $a^b \mod n$. First, let d := 1 and obtain the binary representation of b, say (b_1, b_2, \ldots, b_k) where each $b_i \in \mathbf{Z}/2\mathbf{Z}$. For $i := 1, \ldots, k$, let $d := d^2 \mod n$ and if $b_i = 1$ then let $d := da \mod n$. This algorithm is implemented in the function power_mod. We now use the function power_mod to encrypt our message:

```
sage: m = 72697676798779827668
sage: e = 1850567623300615966303954877
sage: n = 4951760154835678088235319297
sage: c = power_mod(m, e, n); c
630913632577520058415521090
```

Thus c = 630913632577520058415521090 is the ciphertext. To recover our plaintext, we raise c to the power of d and reduce the result modulo n. Again, we use modular exponentiation via repeated squaring in the decryption process:

```
sage: m = 72697676798779827668
sage: c = 630913632577520058415521090
sage: d = 4460824882019967172592779313
sage: n = 4951760154835678088235319297
sage: power_mod(c, d, n)
72697676798779827668
sage: power_mod(c, d, n) == m
True
```

Notice in the last output that the value 72697676798779827668 is the same as the integer that represents our original message. Hence we have recovered our plaintext.

Acknowledgements

- 1. 2009-07-25: Ron Evans (Department of Mathematics, UCSD) reported a typo in the definition of greatest common divisors. The revised definition incorporates his suggestions.
- 2008-11-04: Martin Albrecht (Information Security Group, Royal Holloway, University of London), John Cremona (Mathematics Institute, University of Warwick) and William Stein (Department of Mathematics, University of Washington) reviewed this tutorial. Many of their invaluable suggestions have been incorporated into this document.

Bibliography

12.1.8 Sage and Coding Theory

Author: David Joyner and Robert Miller (2008), edited by Ralf Stephan

This brief paper surveys recent work in Sage on implementing algorithms to compute with linear block codes.

Included in Sage is the group theory package GAP [GAP] and GUAVA [GUAVA], GAP's coding theory package. All of GUAVA's functions can be accessed within Sage.

General coding theory functions

Many of the following coding theory functions are pure Python and do not call GUAVA:

1. LinearCode class definition; LinearCodeFromVectorspace conversion function.

```
[1, 0, 0, 7, 7, 0, 0, 1]
sage: C.weight_distribution()
[1, 0, 0, 7, 7, 0, 0, 1]
```

This function also enables you to create your own code functions. The following function implements the hexacode.

2. The spectrum (weight distribution), minimum_distance programs (calling Steve Linton's C programs in GAP), characteristic_function (as in [vanLint]), and several implementations of the Duursma zeta function (zeta_polynomial, zeta_function, chinen_polynomial, for example),

```
sage: C = codes.HammingCode(3,GF(2))
sage: C.zeta_polynomial()
2/5*T^2 + 2/5*T + 1/5
sage: C = best_known_linear_code(6,3,GF(2)) # known bug
sage: C.minimum_distance() # known bug
3
sage: C.zeta_polynomial() # known bug
2/5*T^2 + 2/5*T + 1/5
```

3. gen_mat, check_mat, decode, dual_code, extended_code, binomial_moment for LinearCode.

The i-th binomial moment of the $[n, k, d]_q$ -code C is

$$B_i(C) = \sum_{S, |S|=i} \frac{q^{k_S} - 1}{q - 1}$$

where k_S is the dimension of the shortened code C_{J-S} , J = [1, 2, ..., n].

```
sage: C = codes.HammingCode(3,GF(2))
sage: C.binomial_moment(2)
0
sage: C.binomial_moment(3) # known bug
0
sage: C.binomial_moment(4) # long time
35
sage: C = codes.HammingCode(3,GF(2))
```

```
sage: MS = MatrixSpace(GF(2),1,7)
sage: F = GF(2); a = F.gen()
sage: v1 = [a,a,F(0),a,a,F(0),a]
sage: C.decode(v1) # known bug
(1, 0, 0, 1, 1, 0, 1)
```

Decoding, at the moment, merely uses syndrome decoding via GUAVA.

- 4. Boolean-valued functions such as "==", is_self_dual, is_self_orthogonal, is_permutation_automorphism,
- 5. permutation methods: automorphism_group_binary_code, is_permutation_automorphism, standard_form, module_composition_factors.

This latter function simply calls up the MeatAxe record from GAP.

```
sage: C = codes.HammingCode(3,GF(2))
sage: G = C.automorphism_group_binary_code(); G # known bug
Permutation Group with generators [(2,3)(5,7), (2,5)(3,7),
                   (2,3,7,5)(4,6),(2,4)(6,7),(1,2)(3,4)
sage: G.order() # known bug
sage: C = codes.HammingCode(3,GF(2))
sage: C.generator_matrix() # known bug
[1 0 0 1 0 1 0]
[0 1 0 1 0 1 1]
[0 0 1 1 0 0 1]
[0 \ 0 \ 0 \ 0 \ 1 \ 1 \ 1]
sage: C.redundancy_matrix() # known bug
[1 1 0]
[1 1 1]
[1 0 1]
[0 1 1]
sage: C.standard_form()[0].generator_matrix() # known bug
[1 0 0 0 1 1 0]
[0 1 0 0 1 1 1]
[0 0 1 0 1 0 1]
[0 \ 0 \ 0 \ 1 \ 0 \ 1 \ 1]
sage: MS = MatrixSpace(GF(2), 4, 8)
sage: G = MS([[1,0,0,0,1,1,1,1,0],[0,1,1,1,0,0,0,0]), # known bug
               [0,0,0,0,0,0,0,1],[0,0,0,0,0,1,0,0]]) # known bug
. . . . :
sage: C = codes.LinearCode(G)
                                                      # known bug
sage: gp = C.automorphism_group_binary_code()
                                                     # known bua
sage: C.module_composition_factors(gp)
                                                      # known bug
[ rec(
  field := GF(2),
  isMTXModule := true,
  dimension := 1,
  generators := [ [ [Z(2)^0] ], [ [Z(2)^0] ], [ [Z(2)^0] ],
      [ [ Z(2)^0 ] ], [ [ Z(2)^0 ] ], [ [ Z(2)^0 ] ] ],
  smashMeataxe := rec(
      algebraElement :=
       [ [ [ 5, 3 ], [ 5, 3 ] ], [ Z(2)^0, Z(2)^0, 0*Z(2), Z(2)^0,
              0*Z(2), Z(2)^0, Z(2)^0, Z(2)^0, Z(2)^0
      algebraElementMatrix := [0 * Z(2)],
      characteristicPolynomial := x_1,
      charpolFactors := x_1,
      nullspaceVector := [Z(2)^0],
```

```
ndimFlag := 1 ),
IsIrreducible := true ), rec(
field := GF(2),
isMTXModule := true,
dimension := 1,
generators := [ [ [Z(2)^0] ], [ [Z(2)^0] ], [ [Z(2)^0] ],
        [ [ Z(2)^0 ] ], [ [ Z(2)^0 ] ], [ [ Z(2)^0 ] ] ],
smashMeataxe := rec(
        algebraElement :=
          [ [ [ 5, 2 ], [ 1, 2 ] ], [ 0*Z(2), 0*Z(2), 0*Z(2), 0*Z(2),
                         Z(2)^0, 0*Z(2), Z(2)^0, 0*Z(2)],
        algebraElementMatrix := [ [0 \times Z(2)] ],
        characteristicPolynomial := x_1,
        charpolFactors := x_1,
        nullspaceVector := [Z(2)^0],
        ndimFlag := 1),
IsIrreducible := true ), rec(
field := GF(2),
isMTXModule := true,
dimension := 1,
generators := [ [ Z(2)^0 ] ], [ Z(2)^0 ] ], [ Z(2)^0 ] ],
         [ [ Z(2)^0 ] ], [ [ Z(2)^0 ] ], [ [ Z(2)^0 ] ] ],
smashMeataxe := rec(
        algebraElement :=
          [ [ [ 4, 2 ], [ 7, 4 ] ], [ 0*Z(2), Z(2)^0, Z(2)^0, 0*Z(2),
                         Z(2)^0, Z(2)
        algebraElementMatrix := [ [ 0 \times Z(2) ] ],
        characteristicPolynomial := x_1,
        charpolFactors := x_1,
        nullspaceVector := [Z(2)^0],
        ndimFlag := 1 ),
IsIrreducible := true ), rec(
field := GF(2),
isMTXModule := true,
dimension := 1,
generators := [ [ Z(2)^0 ] ], [ [ Z(2)^0 ] ], [ [ Z(2)^0 ] ],
        [ [ Z(2)^0 ] ], [ [ Z(2)^0 ] ], [ [ Z(2)^0 ] ] ],
smashMeataxe := rec(
        algebraElement :=
          [ [ [ 4, 6 ], [ 1, 6 ] ], [ 0*Z(2), Z(2)^0, Z(2)^0, 0*Z(2),
                         Z(2)^0, 0*Z(2), Z(2)^0, Z(2)^0, [],
        algebraElementMatrix := [ [ Z(2)^0 ] ],
        characteristicPolynomial := x_1+Z(2)^0,
        charpolFactors := x_1+Z(2)^0,
        nullspaceVector := [Z(2)^0],
        ndimFlag := 1),
IsIrreducible := true ) ]
```

6. design-theoretic methods: assmus_mattson_designs (implementing the Assmus-Mattson Theorem).

Theorem 1. (Assmus and Mattson Theorem. par. 8.4, page 303 of [HP]) Let $A_0, A_1, ..., A_n$ be the weight distribution of the codewords in a binary linear [n, k, d] code C, and let ${}^1A_0^{\perp}, A_1^{\perp}, ..., A_n^{\perp}$ be the weight distribution of the codewords in its dual $[n, n-k, d^{\perp}]$ code C^{\perp} . Fix a t, 0 < t < d, and let

$$s = |\{i \mid A_i^{\perp} \neq 0, 0 < i \le n - t\}|.$$

Assume $s \leq d - t$.

¹ For typographical reasons, the output of the program assmus_mattson_designs uses C^* instead of C^{\perp} .

- If $A_i \neq 0$ and $d \leq i \leq n$ then $C_i = \{c \in C \mid wt(c) = i\}$ holds a simple t-design.
- If $A_i^{\perp} \neq 0$ and $d^{\perp} \leq i \leq n-t$ then $C_i^{\perp} = \{c \in C * \mid wt(c) = i\}$ holds a simple t-design.

Some of the terms in the above theorem are recalled below (see for details). A **block design** is a pair (X, B), where X is a non-empty finite set of v > 0 elements called **points**, and B is a non-empty finite multiset of size b whose elements are called **blocks**, such that each block is a non-empty finite multiset of k points. A design without repeated blocks is called a **simple** block design. If every subset of points of size t is contained in exactly λ blocks the the block design is called a $t - (v, k, \lambda)$ design (or simply a t-design when the parameters are not specified). When $\lambda = 1$ then the block design is called a S(t, k, v) Steiner system.

In the Assmus and Mattson Theorem, X is the set $\{1, 2, ..., n\}$ of coordinate locations and $B = \{supp(c) \mid c \in C_i\}$ is the set of supports of the codewords of C of weight i. Therefore, the parameters of the t-design for C_i are

$$\begin{split} t &= \text{given}, \\ v &= n, \\ k &= i, \quad \text{(this k is not to be confused with $\dim(C)$!),} \\ b &= A_i, \\ \lambda &= b * \frac{\binom{k}{t}}{\binom{v}{t}} \end{split}$$

(by Theorem 8.1.6, p. 294, in [HP]).

Setting the mode="verbose" option prints out the values of the parameters.

The first example below means that the binary [24,12,8]-code C has the property that the (support of the) codewords of weight 8 (resp, 12, 16) form a 5-design. Similarly for its dual code C^{\perp} (of course $C = C^{\perp}$ in this case, so this info is extraneous). The test fails to produce 6-designs (ie, the hypotheses of the theorem fail to hold, not that the 6-designs definitely don't exist). The command assmus_mattson_designs(C,5,mode="verbose") returns the same value but prints out more detailed information.

The second example below illustrates the blocks of the 5-(24, 8, 1) design (i.e., the S(5, 8, 24) Steiner system).

EXAMPLES:

```
sage: C = codes.ExtendedBinaryGolayCode() # example 1
sage: C.assmus_mattson_designs(5)
['weights from C: ',
[8, 12, 16, 24],
'designs from C: ',
[[5, (24, 8, 1)], [5, (24, 12, 48)], [5, (24, 16, 78)], [5, (24, 24, 1)]],
'weights from C*: ',
[8, 12, 16],
'designs from C*: ',
[[5, (24, 8, 1)], [5, (24, 12, 48)], [5, (24, 16, 78)]]]
sage: C.assmus_mattson_designs(6)
0
sage: X = range(24) # example 2
sage: blocks = [c.support() for c in C if hamming_weight(c) == 8] # known bug
sage: len(blocks)
# known bug
```

The method automorphism_group_binary_code is actually an interface to an extremely fast implementation written by the second author. It uses an open-source implementation of permutation backtracking, written by Robert Miller and developed into a Sage module called NICE. This package is described more fully in [Miller1].

A permutation $g \in S_n$ of the fixed basis gives rise to a permutation of the vectors, or words, in $GF(2)^n$, sending (w_i) to $(w_{g(i)})$. The (**permutation**) automorphism group of the code C is the set of permutations of the indices that

bijectively map C to itself. Sage uses a partition refinement algorithm to compute the automorphism group of any binary code. In future work, this will be extended to other base rings.

Native constructions

Sage contains GUAVA but most of GUAVA's functions have not been implemented in Python, so they must be called via the GAP interface. (See the GUAVA manual: https://code.google.com/p/guava-libraries/ for details on the syntax of GUAVA.)

In addition, here are some of the special codes implemented natively in Python:

• BCHCode - A 'Bose-Chaudhuri-Hockenghem code' (or BCH code, for short) is the largest possible cyclic code of length n over field F = GF(q), whose generator polynomial has zeros (contained in) $\{\alpha^b, \alpha^{b+1}, \dots \alpha^{b+\delta-2}\}$, where α is a primitive n^{th} root of unity in the splitting field $GF(q^m)$, b is an integer $0 \le b \le n - \delta + 1$ and m is the multiplicative order of q modulo n.

SEEALSO: Wikipedia article BCH_code

EXAMPLES:

```
sage: FF.<a> = GF(3^2,"a")
sage: x = PolynomialRing(FF,"x").gen()
sage: L = [b.minpoly() for b in [a,a^2,a^3]]; g = LCM(L)
sage: f = x^(8)-1
sage: g.divides(f)
True
sage: C = codes.CyclicCode(8,g); C
Linear code of length 8, dimension 4 over Finite Field of size 3
sage: C.minimum_distance()
4
sage: C = codes.BCHCode(8,3,GF(3),1); C
Linear code of length 8, dimension 4 over Finite Field of size 3
sage: C.minimum_distance()
4
sage: C = codes.BCHCode(8,5,GF(3)); C
Linear code of length 8, dimension 3 over Finite Field of size 3
sage: C.minimum_distance()
5
```

• BinaryGolayCode, ExtendedBinaryGolayCode, TernaryGolayCode, - the well-known "extremal" Golay codes: Wikipedia article Golay_code

EXAMPLES:

```
sage: C = codes.ExtendedBinaryGolayCode()
sage: C
Linear code of length 24, dimension 12 over Finite Field of size 2
sage: C.minimum_distance()
8
sage: C.is_self_dual()
True
sage: C = codes.TernaryGolayCode()
sage: C
Linear code of length 11, dimension 6 over Finite Field of size 3
sage: C.minimum_distance()
```

• Cyclic codes - CyclicCodeFromGeneratingPolynomial (= CyclicCode), CyclicCodeFromCheckPolynomial: Wikipedia article Cyclic_code

EXAMPLES:

```
sage: P.\langle x \rangle = PolynomialRing(GF(3), "x")
sage: q = x-1
sage: C = codes.CyclicCodeFromGeneratingPolynomial(4,q); C
Linear code of length 4, dimension 3 over Finite Field of size 3
sage: P.<x> = PolynomialRing(GF(4, "a"), "x")
sage: q = x^3+1
sage: C = codes.CyclicCodeFromGeneratingPolynomial(9,g); C
Linear code of length 9, dimension 6 over Finite Field in a of size 2^2
sage: P.<x> = PolynomialRing(GF(2), "x")
sage: g = x^3+x+1
sage: C = codes.CyclicCodeFromGeneratingPolynomial(7,g); C
Linear code of length 7, dimension 4 over Finite Field of size 2
sage: C.generator_matrix()
[1 1 0 1 0 0 0]
[0 1 1 0 1 0 0]
[0 0 1 1 0 1 0]
[0 0 0 1 1 0 1]
sage: g = x+1
sage: C = codes.CyclicCodeFromGeneratingPolynomial(4,q); C
Linear code of length 4, dimension 3 over Finite Field of size 2
sage: C.generator_matrix()
[1 1 0 0]
[0 1 1 0]
[0 0 1 1]
sage: P.<x> = PolynomialRing(GF(3),"x")
sage: C = codes.CyclicCodeFromCheckPolynomial(4,x + 1); C
Linear code of length 4, dimension 1 over Finite Field of size 3
sage: C = codes.CyclicCodeFromCheckPolynomial(4, x^3 + x^2 + x + 1); C
Linear code of length 4, dimension 3 over Finite Field of size 3
sage: C.generator_matrix()
[2 1 0 0]
[0 2 1 0]
[0 0 2 1]
```

• DuadicCodeEvenPair, DuadicCodeOddPair - Constructs the "even" (resp. "odd") pair of duadic codes associated to a "splitting" S_1 , S_2 of n. This is a special type of cyclic code whose generator is determined by S_1 , S_2 . See chapter 6 in [HP].

EXAMPLES:

```
sage: from sage.coding.code_constructions import is_a_splitting
sage: n = 11; q = 3
sage: C = Zmod(n).cyclotomic_cosets(q); C
[[0], [1, 3, 4, 5, 9], [2, 6, 7, 8, 10]]
sage: S1 = C[1]
sage: S2 = C[2]
sage: is_a_splitting(S1,S2,11)
True
sage: codes.DuadicCodeOddPair(GF(q),S1,S2)
(Linear code of length 11, dimension 6 over Finite Field of size 3, Linear code of length 11, dimension 6 over Finite Field of size 3)
```

This is consistent with Theorem 6.1.3 in [HP].

• HammingCode - the well-known Hamming code.

The r^{th} Hamming code over F = GF(q) is an [n, k, d] code with length $n = (q^r - 1)/(q - 1)$, dimension $k = (q^r - 1)/(q - 1) - r$ and minimum distance d = 3. The parity check matrix of a Hamming code has rows

consisting of all nonzero vectors of length r in its columns, modulo a scalar factor so no parallel columns arise. A Hamming code is a single error-correcting code.

SEEALSO: Wikipedia article Hamming_code

EXAMPLES:

```
sage: codes.HammingCode(3,GF(2))
Linear code of length 7, dimension 4 over Finite Field of size 2
sage: C = codes.HammingCode(3,GF(3)); C
Linear code of length 13, dimension 10 over Finite Field of size 3
sage: C.minimum_distance()
3
sage: C = codes.HammingCode(3,GF(4,'a')); C
Linear code of length 21, dimension 18 over Finite Field in a of size 2^2
```

 LinearCodeFromCheckMatrix - for specifing the code using the check matrix instead of the generator matrix.

A linear [n, k]-code C is uniquely determined by its generator matrix G and check matrix H. These objects and morphisms fit into the following short exact sequence,

$$0 \to \mathbf{F}^k \stackrel{G}{\to} \mathbf{F}^n \stackrel{H}{\to} \mathbf{F}^{n-k} \to 0.$$

Here, "short exact" means (a) the arrow G is injective, i.e., G is a full-rank $k \times n$ matrix, (b) the arrow H is surjective, and (c) image(G) = kernel(H).

EXAMPLES:

```
sage: C = codes.HammingCode(3,GF(2))
sage: H = C.parity_check_matrix(); H
                                        # known bug
[1 0 0 1 1 0 1]
[0 1 0 1 0 1 1]
[0 0 1 1 1 1 0]
sage: codes.LinearCodeFromCheckMatrix(H) == C # known bug
sage: C = codes.HammingCode(2,GF(3))
sage: H = C.parity_check_matrix(); H
                                                         # known bug
[1 0 2 2]
[0 1 2 1]
sage: codes.LinearCodeFromCheckMatrix(H) == C # known bug
True
sage: C = codes.RandomLinearCode(10,5,GF(4,"a"))
sage: H = C.parity_check_matrix()
sage: codes.LinearCodeFromCheckMatrix(H) == C # known bug
True
```

• QuadraticResidueCodeEvenPair, QuadraticResidueCodeOddPair: Quadratic residue codes of a given odd prime length and base ring either don't exist at all or occur as 4-tuples - a pair of "odd-like" codes and a pair of "even-like" codes. If n>2 is prime then (Theorem 6.6.2 in [HP]) a QR code exists over GF(q) if and only if q is a quadratic residue \pmod{n} . Here they are constructed as "even-like" (resp., "odd-like") duadic codes associated the splitting $(Q,N)\pmod{n}$, where Q is the set of non-zero quadratic residues and N is the non-residues.

QuadraticResidueCode (a special case) and ExtendedQuadraticResidueCode are included as well.

```
sage: codes.QuadraticResidueCodeEvenPair(17,GF(13))
(Linear code of length 17, dimension 8 over Finite Field of size 13,
```

```
Linear code of length 17, dimension 8 over Finite Field of size 13)

sage: codes.QuadraticResidueCodeEvenPair(17,GF(2))

(Linear code of length 17, dimension 8 over Finite Field of size 2,
   Linear code of length 17, dimension 8 over Finite Field of size 2)

sage: codes.QuadraticResidueCodeEvenPair(13,GF(9,"z"))

(Linear code of length 13, dimension 6 over Finite Field in z of size 3^2,
   Linear code of length 13, dimension 6 over Finite Field in z of size 3^2,
   Linear code of length 13, dimension 6 over Finite Field in z of size 3^2)

sage: C1 = codes.QuadraticResidueCodeEvenPair(7,GF(2))[0]

sage: C1.is_self_orthogonal()

True

sage: C2 = codes.QuadraticResidueCodeEvenPair(7,GF(2))[1]

sage: C3 = codes.QuadraticResidueCodeOddPair(17,GF(2))[1]

sage: C4 = codes.QuadraticResidueCodeEvenPair(17,GF(2))[1]

sage: C3 == C4.dual_code()

True
```

This is consistent with Theorem 6.6.9 and Exercise 365 in [HP].

- RandomLinearCode Repeatedly applies Sage's random_element applied to the ambient MatrixSpace of the generator matrix until a full rank matrix is found.
- GeneralizedReedSolomonCode Given a finite field \mathbb{F} of order q, let n and k be chosen such that $1 \leq k \leq n \leq q$. Pick n distinct elements of \mathbb{F} , denoted $\{x_1, x_2, ..., x_n\}$. Then, the codewords are obtained by evaluating every polynomial in $\mathbb{F}[x]$ of degree less than k at each x_i :

$$C = \{ (f(x_1), f(x_2), ..., f(x_n)) \mid f \in \mathbb{F}[x], \deg(f) < k \}.$$

C is a [n, k, n - k + 1] code. (In particular, C is MDS ².)

INPUT:

- evaluation_points A list of evaluation points in a finite field F
- dimension The dimension of the code
- column_multipliers (default: None) List of column multipliers in F for this code. All column multipliers are set to 1 if default value is kept.

EXAMPLES:

```
sage: C = codes.GeneralizedReedSolomonCode(GF(7).list()[:6], 4); C
[6, 4, 3] Generalized Reed-Solomon Code over Finite Field of size 7
sage: C.minimum_distance()
3
sage: F.<a> = GF(3^2,"a")
sage: pts = [0,1,a,a^2,2*a,2*a+1]
sage: len(Set(pts)) == 6 # to make sure there are no duplicates
True
sage: C = codes.GeneralizedReedSolomonCode(pts, 4); C
[6, 4, 3] Generalized Reed-Solomon Code over Finite Field in a of size 3^2
sage: C.minimum_distance()
```

• ToricCode - Let P denote a list of lattice points in \mathbb{Z}^d and let T denote a listing of all points in $(\mathbb{F}^x)^d$. Put n=|T| and let k denote the dimension of the vector space of functions $V=Span\{x^e\mid e\in P\}$. The associated toric code C is the evaluation code which is the image of the evaluation map $eval_T:V\to\mathbb{F}^n$, where x^e is the multi-index notation.

² A code C whose parameters satisfy k + d = n + 1 is called **maximum distance separable** or **MDS**.

EXAMPLES:

```
sage: C = codes.ToricCode([[0,0],[1,0],[2,0],[0,1],[1,1]],GF(7))
sage: C
Linear code of length 36, dimension 5 over Finite Field of size 7
sage: C.minimum_distance()
24
sage: P = [ [0,0],[1,1],[1,2],[1,3],[1,4],[2,1],[2,2],[2,3],[3,1],[3,2],[4,1]]
sage: C = codes.ToricCode(P, GF(8,"a"))
sage: C
Linear code of length 49, dimension 11 over Finite Field in a of size 2^3
```

This is in fact a [49, 11, 28] code over GF(8). If you type next C.minimum_distance() and wait overnight (!), you will get 28.

ullet WalshCode - a binary linear $[2^m,m,2^{m-1}]$ code related to Hadamard matrices. Wikipedia article Walsh_code

EXAMPLES:

```
sage: C = codes.WalshCode(4); C
Linear code of length 16, dimension 4 over Finite Field of size 2
sage: C.minimum_distance()
```

Bounds

Regarding bounds on coding theory parameters, this module implements:

• best_known_linear_code_www (interface with codetables.de since A. Brouwer's online tables have been disabled). Explains the construction of the best known linear code over GF(q) with length n and dimension k, courtesy of the www page http://www.codetables.de/.

INPUT:

- n integer, the length of the code
- k integer, the dimension of the code
- F finite field, whose field order must be in [2, 3, 4, 5, 7, 8, 9]
- verbose bool (default=False), print verbose mesSage

EXAMPLES:

• bounds_minimum_distance which call tables in GUAVA (updated May 2006) created by Cen Tjhai instead of the online internet tables. It simply returns the GAP record for that code:

```
sage: print bounds_minimum_distance(10,5,GF(2)) # known bug
rec(
   n := 10,
```

```
k := 5,
q := 2,
references := rec(
    ),
construction :=
 [ <Operation "ShortenedCode">, [ [ <Operation "UUVCode">, [ [
                    <Operation "DualCode">,
                    [ [ <Operation "RepetitionCode">, [ 8, 2 ] ] ],
                [ <Operation "UUVCode">,
                    [ [ <Operation "DualCode">,
                    [ [ <Operation "RepetitionCode">, [ 4, 2 ] ] ],
                      [ <Operation "RepetitionCode">, [ 4, 2 ] ] ] ],
                      [1, 2, 3, 4, 5, 6]]],
lowerBound := 4,
lowerBoundExplanation :=
 [ "Lb(10,5)=4, by shortening of:",
   "Lb(16,11)=4, by the u|u+v construction applied to C1 [8,7,2] and C2 [8,4,4]: ",
    "Lb(8,7)=2, dual of the repetition code",
    "Lb(8,4)=4, by the u|u+v construction applied to C1 [4,3,2] and C2 [4,1,4]: ",
    "Lb(4,3)=2, dual of the repetition code", "Lb(4,1)=4, repetition code"
   ],
upperBound := 4,
upperBoundExplanation := ["Ub(10,5)=4, by the Griesmer bound"])
```

• codesize_upper_bound (n, d, q), for the best known (as of May, 2006) upper bound A(n, d) for the size of a code of length n, minimum distance d over a field of size q.

EXAMPLES:

```
sage: codesize_upper_bound(10, 3, 2) # known bug
85
```

This means that there is a (10, 85, 3) binary (non-linear) code. Since $85 > 2^6$, this is a better code that a [10, 6, 3] binary (linear) code, assuming one exists. Let's use best_known_linear_code_www to find out:

```
sage: L = best_known_linear_code_www(10, 6, GF(2)) # known bug
sage: print L
                                                   # known bug
Construction of a linear code
[10,6,3] over GF(2):
[1]: [4, 1, 4] Cyclic Linear Code over GF(2)
     RepetitionCode of length 4
[2]: [4, 3, 2] Cyclic Linear Code over GF(2)
    Dual of the RepetitionCode of length 4
[3]: [8, 4, 4] Quasicyclic of degree 2 Linear Code over GF(2)
    PlotkinSum of [2] and [1]
[4]: [8, 7, 2] Cyclic Linear Code over GF(2)
    Dual of the RepetitionCode of length 8
[5]: [16, 11, 4] Linear Code over GF(2)
    PlotkinSum of [4] and [3]
[6]: [15, 11, 3] Linear Code over GF(2)
    Puncturing of [5] at 1
[7]: [10, 6, 3] Linear Code over GF(2)
     Shortening of [6] at { 11 .. 15 }
last modified: 2001-01-30
```

Not only does a [10, 6, 3] binary linear code exist, the value d = 3 is the minimum distance is best known for n = 10, k = 6.

• dimension_upper_bound (n, d, q), an upper bound $B(n,d) = B_q(n,d)$ for the dimension of a linear

code of length n, minimum distance d over a field of size q.

EXAMPLES:

sage: codes.bounds.dimension_upper_bound(10, 3, 2)
6

This was established in the example above.

- gilbert_lower_bound (n, q, d), a lower bound for number of elements in the largest code of minimum distance d in \mathbb{F}_q^n .
- gv_info_rate(n,delta,q), namely $log_q(GLB)/n$, where GLB is the Gilbert lower bound above and delta = d/n.

Let

$$R = R(C) = \frac{k}{n},$$

which measures the information rate of the code, and

$$\delta = \delta(C) = \frac{d}{n},$$

which measures the error correcting ability of the code. Let Σ_q denote the set of all $(\delta, R) \in [0, 1]^2$ such that there exists a sequence C_i , i = 1, 2, ..., of $[n_i, k_i, d_i]$ -codes for which $\lim_{i \to \infty} d_i/n_1 = \delta$ and $\lim_{i \to \infty} k_i/n_i = R$.

The following theorem describes information-theoretical limits on how "good" a linear code can be.

Theorem 2 (Manin [SS], chapter 1). There exists a continuous decreasing function

$$\alpha_q: [0,1] \to [0,1],$$

such that

- α_q is strictly decreasing on $[0, \frac{q-1}{q}]$,
- $\alpha_q(0) = 1,$
- if $\frac{q-1}{q} \le x \le 1$ then $\alpha_q(x) = 0$,
- $\Sigma_q = \{ (\delta, R) \in [0, 1]^2 \mid 0 \le R \le \alpha_q(\delta) \}.$

Not a single value of $\alpha_q(x)$ is known for $0 < x < \frac{q-1}{q}!$ It is not known whether or not the maximum value of the bound, $R = \alpha_q(\delta)$ is attained by a sequence of linear codes. It is not known whether or not $\alpha_q(x)$ is differentiable for $0 < x < \frac{q-1}{q}$, nor is it known if $\alpha_q(x)$ is convex on $0 < x < \frac{q-1}{q}$. However, the following estimate is known.

Theorem 3 (Gilbert-Varshamov, [SS] chapter 1). We have

$$\alpha_q(x) \ge 1 - x \log_q(q - 1) - x \log_q(x) - (1 - x) \log_q(1 - x).$$

In other words, for each fixed $\epsilon > 0$, there exists an (n, k, d)-code C (which may depend on ϵ) with

$$\begin{split} R(C) + \delta(C) \geq & 1 - \delta(C) \log_q(\frac{q-1}{q}) - \delta(C) \log_q(\delta(C)) \\ & - (1 - \delta(C)) \log_q(1 - \delta(C)) - \epsilon. \end{split}$$

The curve $(\delta, 1 - \delta \log_q(\frac{q-1}{q}) - \delta \log_q(\delta) - (1-\delta) \log_q(1-\delta))$ is called the **Gilbert-Varshamov curve**.

• gv_bound_asymp (delta, q), asymptotic analog of the Gilbert lower bound.

```
Sage : f = lambda x: gv_bound_asymp(x,2)
Sage : plot(f,0,1/2)
```

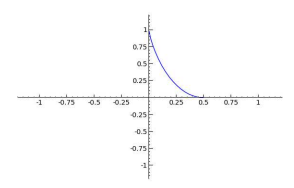


Figure 12.7: Figure 1: Plot of the Gilbert-Varshamov curve using Sage (i.e., y = gv_bound_asymp(x, 2)).

- plotkin_upper_bound(n,q,d)
- plotkin_bound_asymp (delta, q), asymptotic analog of the Plotkin upper bound.

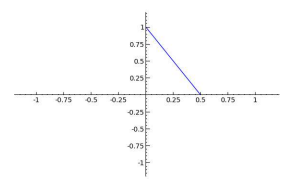


Figure 12.8: Figure 2: Plot using Sage of $y = plotkin_bound_asymp(x, 2)$.

- griesmer_upper_bound (n, q, d), the Griesmer upper bound.
- elias_upper_bound(n,q,d), the Elias upper bound.
- elias_bound_asymp (delta, q), asymptotic analog of the Elias upper bound.
- hamming_upper_bound(n,q,d), the Hamming upper bound.
- \bullet hamming_bound_asymp (delta, q), asymptotic analog of the Hamming upper bound.
- singleton_upper_bound(n,q,d), the Singleton upper bound.
- singleton_bound_asymp (delta, q), asymptotic analog of the Singleton upper bound.
- mrrw1_bound_asymp (delta,q), "first" asymptotic McEliese-Rumsey-Rodemich-Welsh upper bound for the information rate.

Here are all the bounds together:

```
sage: f1 = lambda x: codes.bounds.gv_bound_asymp(x,2)
sage: P1 = plot(f1,0,1/2,linestyle=":")
sage: f2 = lambda x: codes.bounds.plotkin_bound_asymp(x,2)
sage: P2 = plot(f2,0,1/2,linestyle="--")
```

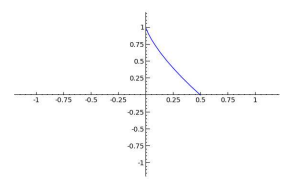


Figure 12.9: Figure 3: Plot using Sage of $y = elias_bound_asymp(x, 2)$.

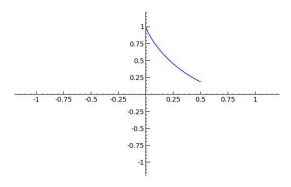


Figure 12.10: Figure 4: Plot using Sage of $y = hamming_bound_asymp(x, 2)$.

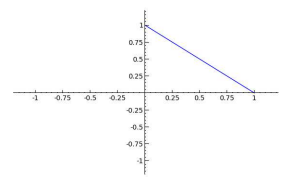


Figure 12.11: Figure 5: Plot using Sage of $y = singleton_bound_asymp(x, 2)$.

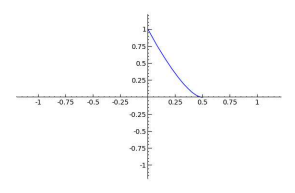


Figure 12.12: Figure 6: Plot using Sage of $y = mrrw1_bound_asymp(x, 2)$.

```
sage: f3 = lambda x: codes.bounds.elias_bound_asymp(x,2)
sage: P3 = plot(f3,0,1/2,rgbcolor=(1,0,0))
sage: f4 = lambda x: codes.bounds.singleton_bound_asymp(x,2)
sage: P4 = plot(f4,0,1/2,linestyle="-.")
sage: f5 = lambda x: codes.bounds.mrrw1_bound_asymp(x,2)
sage: P5 = plot(f5,0,1/2,linestyle="steps")
sage: f6 = lambda x: codes.bounds.hamming_bound_asymp(x,2)
sage: P6 = plot(f6,0,1/2,rgbcolor=(0,1,0))
sage: show(P1+P2+P3+P4+P5+P6)
```

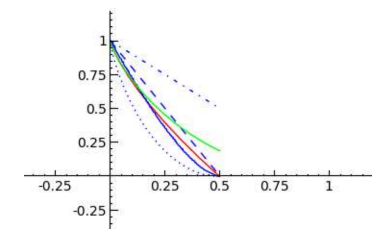


Figure 12.13: Figure 7: Plot of the Gilbert-Varshamov (dotted), Elias (red), Plotkin (dashed), Singleton (dash-dotted), Hamming (green), and MRRW (stepped) curves using Sage.

Self-dual codes

Sage also includes a database of all self-dual binary codes of length ≤ 20 (and some of length 22). The main function is self_dual_codes_binary, which is a case-by-case list of entries, each represented by a Python dictionary.

Format of each entry: a Python dictionary with keys order autgp, spectrum, code, Comment, Type, where

- code a self-dual code C of length n, dimension n/2, over GF(2),
- order autgp order of the permutation automorphism group of C,
- Type the type of C (which can be "I" or "II", in the binary case),

- spectrum the spectrum $[A_0, A_1, ..., A_n]$,
- Comment possibly an empty string.

In fact, in Table 9.10 of , the number B_n of inequivalent self-dual binary codes of length n is given:

n	2	4	6	8	10	12	14	16	18	20	22	24	26	28	30
B_n	1	1	1	2	2	3	4	7	9	16	25	55	103	261	731

According to an entry in Sloane's Online Encyclopedia of Integer Sequences, http://oeis.org/A003179, the next 2 entries are: 3295, 24147.

EXAMPLES:

```
sage: C = self_dual_codes_binary(10)["10"]
sage: C["0"]["code"] == C["0"]["code"].dual_code()
True
sage: C["1"]["code"] == C["1"]["code"].dual_code()
True
sage: len(C.keys()) # number of inequiv sd codes of length 10
2
sage: C = self_dual_codes_binary(12)["12"]
sage: C["0"]["code"] == C["0"]["code"].dual_code()
True
sage: C["1"]["code"] == C["1"]["code"].dual_code()
True
sage: C["2"]["code"] == C["2"]["code"].dual_code()
```

These Sage commands simply show that the two inequivalent self-dual binary codes of length 10, and the two inequivalent self-dual binary codes of length 12, are indeed self dual.

A lot of work on the classification of doubly even self-orthogonal codes using Sage can be found at http://www.rlmiller.org/de_codes/.

The number of permutation equivalence classes of all doubly even [n,k]-codes is shown in the table at http://www.rlmiller.org/de_codes/, and the list of codes so far discovered is linked from the list entries. Each link on that webpage points to a Sage object file, which when loaded (e.g., Sage: L = load('24_12_de_codes.sobj')) is a list of matrices in standard form. The algorithm is described in .

REFERENCES:

12.1.9 A Brief Introduction to Polytopes in Sage

Author: sarah-marie belcastro <smbelcas@toroidalsnark.net>

If you already know some convex geometry *a la* Grünbaum or Brøndsted, then you may have itched to get your hands dirty with some polytope calculations. Here is a mini-guide to doing just that.

Basics

First, let's define a polytope as the convex hull of a set of points, i.e. given S we compute P = conv(S):

```
sage: P1 = Polyhedron(vertices = [[-5,2], [4,4], [3,0], [1,0], [2,-4], [-3,-1], [-5,-3]]) sage: P1 A 2-dimensional polyhedron in ZZ^2 defined as the convex hull of 4 vertices
```

Notice that Sage tells you the dimension of the polytope as well as the dimension of the ambient space.

Of course, you want to know what this object looks like:

```
sage: P1.plot()
Graphics object consisting of 6 graphics primitives
```

Even in only 2 dimensions, it's a pain to figure out what the supporting hyperplanes are. Luckily Sage will take care of that for us.

```
sage: for q in P1.Hrepresentation():
....: print q
An inequality (-4, 1) \times + 12 >= 0
An inequality (1, 7) \times + 26 >= 0
An inequality (1, 0) \times + 5 >= 0
An inequality (2, -9) \times + 28 >= 0
```

That notation is not immediately parseable, because seriously, those do not look like equations of lines (or of halfspaces, which is really what they are).

```
(-4, 1) \times + 12 >= 0 really means (-4, 1) \cdot \vec{x} + 12 \ge 0.
```

So... if you want to define a polytope via inequalities, you have to translate each inequality into a vector. For example, $(-4, 1) \cdot \vec{x} + 12 \ge 0$ becomes (12, -4, 1).

```
sage: altP1 = Polyhedron(ieqs=[(12, -4, 1), (26, 1, 7), (5,1,0), (28, 2, -9)])
sage: altP1.plot()
Graphics object consisting of 6 graphics primitives
```

Other information you might want to pull out of Sage about a polytope is the vertex list, which can be done in two ways:

```
sage: for q in P1.Vrepresentation():
....: print q
A vertex at (-5, -3)
A vertex at (-5, 2)
A vertex at (4, 4)
A vertex at (2, -4)

sage: P1.vertices()
(A vertex at (-5, -3), A vertex at (-5, 2), A vertex at (4, 4), A vertex at (2, -4))
```

Polar duals

Surely you want to compute the polar dual:

```
sage: Pldual = Pl.polar()
sage: Pldual
A 2-dimensional polyhedron in QQ^2 defined as the convex hull of 4 vertices
```

Check it out—we started with an integer-lattice polytope and dualized to a rational-lattice polytope. Let's look at that.

```
sage: Pldual.plot()
Graphics object consisting of 6 graphics primitives
sage: Pl.plot() + Pldual.plot()
Graphics object consisting of 12 graphics primitives
```

Oh, yeah, unless the polytope is unit-sphere-sized, the dual will be a very different size. Let's rescale.

```
sage: ((1/4)*P1).plot() + (4*P1dual).plot()
Graphics object consisting of 12 graphics primitives
```

If you think that looks a little bit shady, you're correct. Here is an example that makes the issue a bit clearer.

```
sage: P2 = Polyhedron(vertices = [[-5,0], [-1,1], [-2,0], [1,0], [-2,-1], [-3,-1], [-5,-1]])
sage: P2
A 2-dimensional polyhedron in ZZ^2 defined as the convex hull of 5 vertices
sage: P2dual = P2.polar(); P2dual
A 2-dimensional polyhedron in QQ^2 defined as the convex hull of 5 vertices
sage: P2.plot() + P2dual.plot()
Graphics object consisting of 14 graphics primitives
```

That is clearly not computing what we think of as the polar dual. But look at this...

```
sage: P2.plot() + (-1*P2dual).plot()
Graphics object consisting of 14 graphics primitives
```

Here is what's going on.

If a polytope P is in \mathbb{Z} , then...

- 1. ...the dual is inverted in some way, which is vertically for polygons.
- 2. ...the dual is taken of P itself.
- 3. ...if the origin is not in P, then an error is returned.

However, if a polytope is *not* in \mathbb{Z} , for example if it's in \mathbb{Q} or RDF, then...

- (1') ...the dual is not inverted.
- (2') ...the dual is taken of P-translated-so-barycenter-is-at-origin.

Keep all of this in mind as you take polar duals.

Polytope Constructions

Minkowski sums! Now with two syntaxes!

```
sage: P1+P2
A 2-dimensional polyhedron in ZZ^2 defined as the convex hull of 8 vertices
sage: P1.Minkowski_sum(P2)
A 2-dimensional polyhedron in ZZ^2 defined as the convex hull of 8 vertices
```

Okay, fine. We should have some 3-dimensional examples, at least. (Note that in order to display polytopes effectively you'll need visualization software such as Javaview and Jmol installed.)

```
sage: P3 = Polyhedron(vertices=[(0,0,0), (0,0,1/2), (0,1/2,0), (1/2,0,0), (3/4,1/5,3/2)]); P3
A 3-dimensional polyhedron in QQ^3 defined as the convex hull of 5 vertices
sage: P4 = Polyhedron(vertices=[(-1,1,0),(1,1,0),(-1,0,1), (1,0,1),(0,-1,1),(0,1,1)]); P4
A 3-dimensional polyhedron in ZZ^3 defined as the convex hull of 6 vertices
sage: P3.plot() + P4.plot()
Graphics3d Object
```

```
sage: (P3+P4).plot()
Graphics3d Object
```

We can also find the intersection of two polytopes... and this too has two syntaxes!

```
sage: int12 = P1.intersection(P2*.5); int12.plot()
Graphics object consisting of 7 graphics primitives
sage: int34 = P3 & P4; int34.plot()
Graphics3d Object
```

Should one wish to translate, one can.

```
sage: transP2 = P2.translation([2,1])
sage: P2.plot() + transP2.plot()
Graphics object consisting of 14 graphics primitives
```

Then of course we can take prisms, pyramids, and bipyramids of polytopes...

```
sage: P2.prism().plot()
Graphics3d Object

sage: P1.pyramid().plot()
Graphics3d Object

sage: P2dual.bipyramid().plot()
Graphics3d Object
```

Okay, fine. Yes, Sage has some kinds of polytopes built in. If you type polytopes. and then press TAB after the period, you'll get a list of pre-built polytopes.

```
sage: P5 = polytopes.hypercube(5)
sage: P6 = polytopes.cross_polytope(3)
sage: P7 = polytopes.simplex(7)
```

Let's look at a 4-dimensional polytope.

```
sage: P8 = polytopes.hypercube(4)
sage: P8.plot()
Graphics3d Object
```

We can see it from a different perspective:

```
sage: P8.schlegel_projection([2,5,11,17]).plot()
Graphics3d Object
```

Queries to polytopes

Once you've constructed some polytope, you can ask Sage questions about it.

```
sage: P1.contains([1,0])
True
```

```
sage: P1.interior_contains([3,0])
False
sage: P3.contains([1,0,0])
False
```

Face information can be useful.

```
sage: int34.f_vector()
(1, 8, 12, 6, 1)
```

Well, geometric information might be *more* helpful... Here we are told which of the vertices form each 2-face:

```
sage: int34.faces(2)
(<1,3,4>, <0,1,3,5>, <0,1,2,4,6>, <2,3,4,5,7>, <2,6,7>, <0,5,6,7>)
```

Yeah, that isn't so useful as it is. Let's figure out the vertex and hyperplane representations of the first face in the list.

```
sage: first2faceofint34 = P3.faces(2)[0]
sage: first2faceofint34.ambient_Hrepresentation(); first2faceofint34.vertices()
(An inequality (1, 0, 0) \times + 0 >= 0,)
(A vertex at (0, 0, 0), A vertex at (0, 0, 1/2), A vertex at (0, 1/2, 0))
```

If you want more... Base class for polyhedra is the first place you want to go.

12.1.10 Draw polytopes in LateX using TikZ

Author: Jean-Philippe Labbé <labbe@math.huji.ac.il>

It is sometimes very helpful to draw 3-dimensional polytopes in a paper. TikZ is a very versatile tool to draw in scientific documents and Sage can deal easily with 3-dimensional polytopes. Finally sagetex makes everything work together nicely between Sage, TikZ and LaTeX. Since version 6.3 of Sage, there is a function for (projection of) polytopes to output a TikZ picture of the polytope. This short tutorial shows how it all works.

Instructions

To put an image of a 3d-polytope in LaTeX using TikZ and Sage, simply follow the instructions:

- Install SageTex (optional but recommended!)
- Put \usepackage{tikz} in the preamble of your article
- Open Sage and change the directory to your article's by the command cd /path/to/article
- Input your polytope, called P for example, to Sage
- Visualize the polytope P using the command P.show (aspect ratio=1)
- This will open an interactive viewer named Jmol, in which you can rotate the polytope. Once the wished view angle is found, right click on the image and select *Console*
- In the dialog box click the button *State*
- Scroll up to the line starting with moveto
- It reads something like moveto 0.0 {x y z angle} scale
- Go back to Sage and type Img = P.projection().tikz([x,y,z],angle)

Img now contains a Sage object of type LatexExpr containing the raw TikZ picture of your polytope

Then, you can either copy-paste it to your article by typing Img in Sage or save it to a file, by doing

```
f = open('Img_poly.tex','w')
f.write(Img)
f.close()
```

Then in the pwd (present working directory of sage, the one of your article) you will have a file named Imq_poly.tex containing the tikzpicture of your polytope.

Customization

You can customize the polytope using the following options in the command P.tikz()

- scale: positive number to scale the polytope,
- edge_color: string (default: blue!95!black) representing colors which tikz recognize,
- facet_color: string (default: blue!95!black) representing colors which tikz recognize,
- vertex_color: string (default: green) representing colors which tikz recognize,
- opacity: real number (default: 0.8) between 0 and 1 giving the opacity of the front facets,
- axis: Boolean (default: False) draw the axes at the origin or not.

Examples

Let's say you want to draw the polar dual of the following (nice!) polytope given by the following list of vertices:

```
[[1,0,1],[1,0,0],[1,1,0],[0,0,-1],[0,1,0],[-1,0,0],[0,1,1],[0,0,1],[0,-1,0]]
```

In Sage, you type:

```
 P = Polyhedron(vertices = [[1,0,1],[1,0,0],[1,1,0],[0,0,-1],[0,1,0],[-1,0,0],[0,1,1],[0,0,1],[0,-1,0]]
```

Then, you visualize the polytope by typing P. show (aspect_ratio=1)

When you found a good angle, follow the above procedure to obtain the values [674,108,-731] and angle=112, for example.

```
Img = P.projection().tikz([674,108,-731],112)
```

Or you may want to customize using the command

```
Img = P.projection().tikz([674,108,-731],112,scale=2, edge_color='orange',facet_color='red',vertex_color='red'
```

Further, you may want to edit deeper the style of the polytope, directly inside the tikzpicture. For example, line 6-9 in the tikzpicture reads:

```
back/.style={loosely dotted, thin},
edge/.style={color=orange, thick},
facet/.style={fill=red,fill opacity=0.400000},
vertex/.style={inner sep=1pt,circle,draw=blue!25!black,fill=blue!75!black,thick,anchor=base}]
```

It is also possible to replace it by the following 4 lines (and adding \usetikzlibrary{shapes} in the preamble)

```
back/.style={loosely dashed,line width=2pt},
edge/.style={color=yellow, line width=2pt},
facet/.style={fill=cyan,fill opacity=0.400000},
vertex/.style={inner sep=4pt,star,star points=7,draw=blue!75!white,fill=blue!85!white,thick,anchor=back.
```

Finally, you may want to tweak your picture my adding labels, elements on vertices, edges, facets, etc.

Automatize using SageTex

For this you need to put

\usepackage{sagetex}

in the preamble of your article

There are different ways to use sagetex and you may create your own. Here are some possibilities.

1. You can directly type in a sagestr in the article:

```
\sagestr{(polytopes.permutahedron(4)).projection().tikz([4,5,6],45,scale=0.75, facet_color='red',ver
```

2. You may create the following tex commands

in your preamble and use them with a sagesilent in your article:

```
\begin{sagesilent}
Polytope = polytopes.great_rhombicuboctahedron()
\end{sagesilent}

\polytopeimg{Polytope}{[276,-607,-746]}{102}{1}
\polytopeimgopt{Polytope}{view=[-907,379,183]}{angle=129}{scale=2}{edge_color='red'}{facet_color='ye.
```

Then, run pdflatex, execute Sage on the file article name.sagetex.sage and run pdflatex again.

For more information on SageTeX, see the tutorial http://www.sagemath.org/doc/tutorial/sagetex.html.

12.1.11 Tutorial: Programming in Python and Sage

Author: Florent Hivert <florent.hivert@univ-rouen.fr>, Franco Saliola @saliola@gmail.com>, et al.

This tutorial is an introduction to basic programming in Python and Sage, for readers with elementary notions of programming but not familiar with the Python language. It is far from exhaustive. For a more complete tutorial, have a look at the Python Tutorial. Also Python's documentation and in particular the standard library can be useful.

A more advanced tutorial presents the notions of objects and classes in Python.

Here are further resources to learn Python:

- Learn Python in 10 minutes ou en français Python en 10 minutes
- Dive into Python is a Python book for experienced programmers. Also available in other languages.
- Discover Python is a series of articles published in IBM's developerWorks technical resource center.

Data structures

In Python, *typing is dynamic*; there is no such thing as declaring variables. The function type() returns the type of an object obj. To convert an object to a type typ just write typ(obj) as in int("123"). The command isinstance(ex, typ) returns whether the expression ex is of type typ. Specifically, any value is *an instance of a class* and there is no difference between classes and types.

The symbol = denotes the affectation to a variable; it should not be confused with == which denotes mathematical equality. Inequality is !=.

The standard types are bool, int, list, tuple, set, dict, str.

- The type bool (*booleans*) has two values: True and False. The boolean operators are denoted by their names or, and, not.
- The Python types int and long are used to represent integers of limited size. To handle arbitrary large integers with exact arithmetic, Sage uses its own type named Integer.
- A *list* is a data structure which groups values. It is constructed using brackets as in [1, 3, 4]. The range () function creates integer lists. One can also create lists using *list comprehension*:

```
[ <expr> for <name> in <iterable> (if <condition>) ]
For example:
sage: [ i^2 for i in range(10) if i % 2 == 0 ]
[0, 4, 16, 36, 64]
```

• A *tuple* is very similar to a list; it is constructed using parentheses. The empty tuple is obtained by () or by the constructor tuple. If there is only one element, one has to write (a,). A tuple is *immutable* (one cannot change it) but it is *hashable* (see below). One can also create tuples using comprehensions:

```
sage: tuple(i^2 for i in range(10) if i % 2 == 0)
(0, 4, 16, 36, 64)
```

• A *set* is a data structure which contains values without multiplicities or order. One creates it from a list (or any iterable) with the constructor set. The elements of a set must be hashable:

```
sage: set([2,2,1,4,5])
{1, 2, 4, 5}

sage: set([[1], [2]])
Traceback (most recent call last):
...
TypeError: unhashable type: 'list'
```

• A *dictionary* is an association table, which associates values to keys. Keys must be hashable. One creates dictionaries using the constructor dict, or using the syntax:

```
{key1 : value1, key2 : value2 ...}
For example:
sage: age = {'toto' : 8, 'mom' : 27}; age
{'mom': 27, 'toto': 8}
```

- Quotes (simple ' ' or double " ") enclose *character strings*. One can concatenate them using +.
- For lists, tuples, strings, and dictionaries, the *indexing operator* is written 1[i]. For lists, tuples, and strings one can also uses *slices* as 1[:], 1[:b], 1[a:], or 1[a:b]. Negative indices start from the end.

- The len() function returns the number of elements of a list, a tuple, a set, a string, or a dictionary. One writes x in C to tests whether x is in C.
- Finally there is a special value called None to denote the absence of a value.

Control structures

In Python, there is no keyword for the beginning and the end of an instructions block. Blocks are delimited solely by means of indentation. Most of the time a new block is introduced by: Python has the following control structures:

• Conditional instruction:

• Inside expression exclusively, one can write:

```
<value> if <condition> else <value>
```

• Iterative instructions:

The else block is executed at the end of the loop if the loop is ended normally, that is neither by a break nor an exception.

- In a loop, continue jumps to the next iteration.
- An iterable is an object which can be iterated through. Iterable types include lists, tuples, dictionaries, and strings.
- An error (also called exception) is raised by:

```
raise <ErrorType> [, error message]
```

Usual errors include ValueError and TypeError.

Functions

Note: Python functions vs. mathematical functions

In what follows, we deal with *functions* is the sense of *programming languages*. Mathematical functions, as manipulated in calculus, are handled by Sage in a different way. In particular it doesn't make sense to do mathematical manipulation such as additions or derivations on Python functions.

One defines a function using the keyword def as:

The result of the function is given by the instruction return. Very short functions can be created anonymously using lambda (remark that there is no instruction return here):

```
lambda <arguments>: <expression>
```

Note: Functional programming

Functions are objects as any other objects. One can assign them to variables or return them. For details, see the tutorial on *Functional Programming for Mathematicians*.

Exercises

Lists

Creating Lists I: [Square brackets] Example:

```
sage: L = [3, Permutation([5,1,4,2,3]), 17, 17, 3, 51]
sage: L
[3, [5, 1, 4, 2, 3], 17, 17, 3, 51]
```

Exercise: Create the list [63, 12, -10, ``a'', 12], assign it to the variable L, and print the list.

```
sage: # edit here
```

Exercise: Create the empty list (you will often need to do this).

```
sage: # edit here
```

Creating Lists II: range The range () function provides an easy way to construct a list of integers. Here is the documentation of the range () function:

```
range([start,] stop[, step]) -> list of integers

Return a list containing an arithmetic progression of integers.
range(i, j) returns [i, i+1, i+2, ..., j-1]; start (!) defaults to 0.
When step is given, it specifies the increment (or decrement). For example, range(4) returns [0, 1, 2, 3]. The end point is omitted!
These are exactly the valid indices for a list of 4 elements.
```

Exercise: Use range () to construct the list [1, 2, ..., 50].

```
sage: # edit here
```

Exercise: Use range () to construct the list of *even* numbers between 1 and 100 (including 100).

```
sage: # edit here
```

Exercise: The step argument for the range() command can be negative. Use range to construct the list [10,7,4,1,-2].

sage: # edit here

See also:

- xrange (): returns an iterator rather than building a list.
- srange (): like range but with Sage integers; see below.
- sxrange(): like xrange but with Sage integers.

Creating Lists III: list comprehensions *List comprehensions* provide a concise way to create lists from other lists (or other data types).

Example We already know how to create the list [1, 2, ..., 16]:

```
sage: range(1,17)
[1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16]
```

Using a *list comprehension*, we can now create the list $[1^2, 2^2, 3^2, \dots, 16^2]$ as follows:

```
sage: [i^2 for i in range(1,17)]
[1, 4, 9, 16, 25, 36, 49, 64, 81, 100, 121, 144, 169, 196, 225, 256]
sage: sum([i^2 for i in range(1,17)])
1496
```

Exercise: [Project Euler, Problem 6]

The sum of the squares of the first ten natural numbers is

$$(1^2 + 2^2 + \dots + 10^2) = 385$$

The square of the sum of the first ten natural numbers is

$$(1+2+...+10)^2 = 55^2 = 3025$$

Hence the difference between the sum of the squares of the first ten natural numbers and the square of the sum is

$$3025 - 385 = 2640$$

Find the difference between the sum of the squares of the first one hundred natural numbers and the square of the sum.

```
sage: # edit here
sage: # edit here
sage: # edit here
```

Filtering lists with a list comprehension A list can be *filtered* using a list comprehension.

Example: To create a list of the squares of the prime numbers between 1 and 100, we use a list comprehension as follows.

```
sage: [p^2 for p in [1,2,..,100] if is_prime(p)]
[4, 9, 25, 49, 121, 169, 289, 361, 529, 841, 961, 1369, 1681, 1849, 2209, 2809, 3481, 3721, 4489, 50-
```

Exercise: Use a *list comprehension* to list all the natural numbers below 20 that are multiples of 3 or 5. Hint:

- To get the remainder of 7 divided by 3 use 7%3.
- To test for equality use two equal signs (==); for example, 3 == 7.

```
sage: # edit here
```

Project Euler, Problem 1: Find the sum of all the multiples of 3 or 5 below 1000.

```
sage: # edit here
```

Nested list comprehensions List comprehensions can be nested!

Examples:

Exercise:

1. A Pythagorean triple is a triple (x, y, z) of positive integers satisfying $x^2 + y^2 = z^2$. The Pythagorean triples whose components are at most 10 are:

Using a filtered list comprehension, construct the list of Pythagorean triples whose components are at most 50:

```
sage: # edit here
sage: # edit here
```

2. Project Euler, Problem 9: There exists exactly one Pythagorean triple for which a+b+c=1000. Find the product abc:

```
sage: # edit here
```

sage: # edit here

Accessing individual elements of lists To access an element of the list L, use the syntax L[i], where i is the index of the item.

Exercise:

```
1. Construct the list L = [1, 2, 3, 4, 3, 5, 6]. What is L[3]?
```

```
2. What is L[1]?
```

```
sage: # edit here
3. What is the index of the first element of L?
sage: # edit here
4. What is L[-1]? What is L[-2]?
sage: # edit here
5. What is L.index(2)? What is L.index(3)?
sage: # edit here
```

Modifying lists: changing an element in a list To change the item in position i of a list L:

```
sage: L = ["a", 4, 1, 8]
sage: L
['a', 4, 1, 8]

sage: L[2] = 0
sage: L
['a', 4, 0, 8]
```

Modifying lists: append and extend To append an object to a list:

```
sage: L = ["a", 4, 1, 8]
sage: L
['a', 4, 1, 8]
sage: L.append(17)
sage: L
['a', 4, 1, 8, 17]
```

To extend a list by another list:

```
sage: L1 = [1,2,3]
sage: L2 = [7,8,9,0]
sage: print L1
[1, 2, 3]
sage: print L2
[7, 8, 9, 0]

sage: L1.extend(L2)
sage: L1
[1, 2, 3, 7, 8, 9, 0]
```

Modifying lists: reverse, sort, ...

```
sage: L = [4,2,5,1,3]
sage: L
[4, 2, 5, 1, 3]
```

```
sage: L.reverse()
sage: L
[3, 1, 5, 2, 4]

sage: L.sort()
sage: L
[1, 2, 3, 4, 5]

sage: L = [3,1,6,4]
sage: sorted(L)
[1, 3, 4, 6]

sage: L
[3, 1, 6, 4]
```

Concatenating Lists To concatenate two lists, add them with the operator +. This is not a commutative operation!

```
sage: L1 = [1,2,3]
sage: L2 = [7,8,9,0]
sage: L1 + L2
[1, 2, 3, 7, 8, 9, 0]
```

Slicing Lists You can slice a list using the syntax L[start : stop : step]. This will return a sublist of L.

Exercise: Below are some examples of slicing lists. Try to guess what the output will be before evaluating the cell:

```
sage: L = range(20)
sage: L
[0, 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19]

sage: L[3:15]
[3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14]

sage: L[3:15:2]
[3, 5, 7, 9, 11, 13]

sage: L[15:3:-1]
[15, 14, 13, 12, 11, 10, 9, 8, 7, 6, 5, 4]

sage: L[:4]
[0, 1, 2, 3]

sage: L[:]
[0, 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19]

sage: L[:-1]
[19, 18, 17, 16, 15, 14, 13, 12, 11, 10, 9, 8, 7, 6, 5, 4, 3, 2, 1, 0]
```

Exercise (Advanced): The following function combines a loop with some of the list operations above. What does the function do?

```
sage: def f(number_of_iterations):
....: L = [1]
....: for n in range(2, number_of_iterations):
....: L = [sum(L[:i]) for i in range(n-1, -1, -1)]
....: return numerical_approx(2*L[0]*len(L)/sum(L), digits=50)
sage: # edit here
```

Tuples

A *tuple* is an *immutable* list. That is, it cannot be changed once it is created. This can be useful for code safety and foremost because it makes tuple *hashable*. To create a tuple, use parentheses instead of brackets:

```
sage: t = (3, 5, [3,1], (17,[2,3],17), 4)
sage: t
(3, 5, [3, 1], (17, [2, 3], 17), 4)
```

To create a singleton tuple, a comma is required to resolve the ambiguity:

```
sage: (1)
1
sage: (1,)
(1,)
```

We can create a tuple from a list, and vice-versa.

```
sage: tuple(range(5))
(0, 1, 2, 3, 4)

sage: list(t)
[3, 5, [3, 1], (17, [2, 3], 17), 4]
```

Tuples behave like lists in many respects:

Operation	Syntax for lists	Syntax for tuples
Accessing a letter	list[3]	tuple[3]
Concatenation	list1 + list2	tuple1 + tuple2
Slicing	list[3:17:2]	tuple[3:17:2]
A reversed copy	list[::-1]	tuple[::-1]
Length	len(list)	len(tuple)

Trying to modify a tuple will fail:

```
sage: t = (5, 'a', 6/5)
sage: t
(5, 'a', 6/5)

sage: t[1] = 'b'
Traceback (most recent call last):
...
TypeError: 'tuple' object does not support item assignment
```

Generators

"Tuple-comprehensions" do not exist. Instead, the syntax produces something called a generator. A generator allows you to process a sequence of items one at a time. Each item is created when it is needed, and then forgotten. This can be very efficient if we only need to use each item once.

```
sage: (i^2 for i in range(5))
<generator object <genexpr> at 0x...>

sage: g = (i^2 for i in range(5))
sage: g[0]
Traceback (most recent call last):
...
TypeError: 'generator' object has no attribute '__getitem__'
sage: [x for x in g]
[0, 1, 4, 9, 16]
g is now empty.
sage: [x for x in g]
[]
```

A nice 'pythonic' trick is to use generators as argument of functions. We do *not* need double parentheses for this:

```
sage: sum(i^2 for i in srange(100001)) 33333833350000
```

Dictionaries

A *dictionary* is another built-in data type. Unlike lists, which are indexed by a range of numbers starting at 0, dictionaries are indexed by *keys*, which can be any immutable objects. Strings and numbers can always be keys (because they are immutable). Dictionaries are sometimes called "associative arrays" in other programming languages.

There are several ways to define dictionaries. One method is to use braces, {}, with comma-separated entries given in the form key:value:

```
sage: d = {3:17, 0.5:[4,1,5,2,3], 0:"goo", 3/2 : 17}
sage: d
{0: 'goo', 0.500000000000000: [4, 1, 5, 2, 3], 3/2: 17, 3: 17}
```

A second method is to use the constructor dict which admits a list (or actually any iterable) of 2-tuples (key, value):

```
sage: dd = dict((i,i^2) for i in xrange(10))
sage: dd
{0: 0, 1: 1, 2: 4, 3: 9, 4: 16, 5: 25, 6: 36, 7: 49, 8: 64, 9: 81}
```

Dictionaries behave as lists and tuples for several important operations.

Operation	Syntax for lists	Syntax for dictionaries
Accessing elements	list[3]	D["key"]
Length	len(list)	len(D)
Modifying	L[3] = 17	D["key"] = 17
Deleting items	del L[3]	del D["key"]

```
sage: d[10]='a'
sage: d
{0: 'goo', 0.500000000000000: [4, 1, 5, 2, 3], 3/2: 17, 3: 17, 10: 'a'}
```

A dictionary can have the same value multiple times, but each key must only appear once and must be immutable:

```
sage: d = {3: 14, 4: 14}
sage: d
{3: 14, 4: 14}

sage: d = {3: 13, 3: 14}
sage: d
{3: 14}

sage: d = {[1,2,3] : 12}
Traceback (most recent call last):
...
TypeError: unhashable type: 'list'
```

Another way to add items to a dictionary is with the update() method which updates the dictionary from another dictionary:

```
sage: d = {}
sage: d
{}

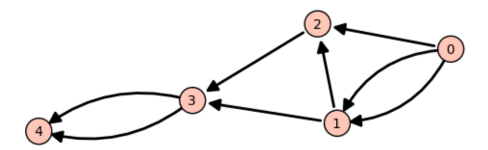
sage: d.update({10 : 'newvalue', 20: 'newervalue', 3: 14, 0.5:[1,2,3]})
sage: d
{0.5000000000000000: [1, 2, 3], 3: 14, 10: 'newvalue', 20: 'newervalue'}
```

We can iterate through the *keys*, or *values*, or both, of a dictionary. Note that, internally, there is no sorting of keys done. In general, the order of keys/values will depend on memory locations can and will differ between different computers and / or repeated runs on the same computer. However, Sage sort the dictionary entries by key when printing the dictionary specifically to make the docstrings more reproducible. However, the Python methods keys() and values() do not sort for you. If you want your output to be reproducable, then you have to sort it first just like in the examples below:

```
sage: d = {10 : 'newvalue', 20: 'newervalue', 3: 14, 0.5:(1,2,3)}
sage: sorted([key for key in d])
[0.50000000000000, 3, 10, 20]
                 # random order
sage: d.keys()
[0.50000000000000, 10, 3, 20]
sage: sorted(d.keys())
[0.50000000000000, 3, 10, 20]
sage: d.values()
                 # random order
[(1, 2, 3), 'newvalue', 14, 'newervalue']
sage: set(d.values()) == set([14, (1, 2, 3), 'newvalue', 'newervalue'])
True
                  # random order
sage: d.items()
[(0.5000000000000, (1, 2, 3)), (10, 'newvalue'), (3, 14), (20, 'newervalue')]
```

```
sage: sorted([(key, value) for key, value in d.items()])
[(0.50000000000000, (1, 2, 3)), (3, 14), (10, 'newvalue'), (20, 'newervalue')]
```

Exercise: Consider the following directed graph.



Create a dictionary whose keys are the vertices of the above directed graph, and whose values are the lists of the vertices that it points to. For instance, the vertex 1 points to the vertices 2 and 3, so the dictionary will look like:

```
d = { ..., 1:[2,3], ... }
sage: # edit here

Then try:
sage: g = DiGraph(d)
sage: g.plot()
```

Using Sage types: The srange command

Example: Construct a 3×3 matrix whose (i,j) entry is the rational number $\frac{i}{j}$. The integers generated by range () are Python int's. As a consequence, dividing them does euclidean division:

```
sage: matrix([[ i/j for j in range(1,4)] for i in range(1,4)])
[1 0 0]
[2 1 0]
[3 1 1]
```

Whereas dividing a Sage Integer by a Sage Integer produces a rational number:

```
sage: matrix([[ i/j for j in srange(1,4)] for i in srange(1,4)])
[ 1 1/2 1/3]
[ 2 1 2/3]
[ 3 3/2 1]
```

Modifying lists has consequences!

Try to predict the results of the following commands:

```
sage: a = [1, 2, 3]
sage: L = [a, a, a]
```

```
[[1, 2, 3], [1, 2, 3], [1, 2, 3]]

sage: a.append(4)
sage: L
[[1, 2, 3, 4], [1, 2, 3, 4], [1, 2, 3, 4]]

Now try these:

sage: a = [1, 2, 3]
sage: L = [a, a, a]
sage: L
[[1, 2, 3], [1, 2, 3], [1, 2, 3]]

sage: a = [1, 2, 3, 4]
sage: L
[[1, 2, 3], [1, 2, 3], [1, 2, 3]]

sage: L
[[1, 2, 3], [1, 2, 3], [1, 2, 3]]
```

sage: L

This is known as the reference effect. You can use the command deepcopy () to avoid this effect:

```
sage: a = [1,2,3]
sage: L = [deepcopy(a), deepcopy(a)]
sage: L
[[1, 2, 3], [1, 2, 3]]

sage: a.append(4)
sage: L
[[1, 2, 3], [1, 2, 3]]
```

The same effect occurs with dictionaries:

```
sage: d = {1:'a', 2:'b', 3:'c'}
sage: dd = d
sage: d.update( { 4:'d' } )
sage: dd
{1: 'a', 2: 'b', 3: 'c', 4: 'd'}
```

Loops and Functions

For more verbose explanation of what's going on here, a good place to look at is the following section of the Python tutorial: http://docs.python.org/tutorial/controlflow.html

While Loops

While loops tend not to be used nearly as much as for loops in Python code:

```
sage: i = 0
sage: while i < 10:
....: print i
....: i += 1</pre>
```

```
0
1
2
3
4
5
6
7
8
9
sage: i = 0
sage: while i < 10:
....: if i % 2 == 1:
          i += 1
...: cont ...: print i ...: i += 1
              continue
0
2
4
6
8
```

Note that the truth value of the clause expression in the *while* loop is evaluated using bool:

```
sage: bool(True)
True
sage: bool('a')
True
sage: bool(1)
True
sage: bool(0)
False
sage: i = 4
sage: while i:
....: print i
         i -= 1
. . . . :
4
3
2
1
```

For Loops

Here is a basic *for* loop iterating over all of the elements in the list 1:

```
sage: l = ['a', 'b', 'c']
sage: for letter in l:
....: print letter
```

b c

The range () function is very useful when you want to generate arithmetic progressions to loop over. Note that the end point is never included:

```
sage: range?
sage: range(4)
[0, 1, 2, 3]
sage: range(1, 5)
[1, 2, 3, 4]
sage: range(1, 11, 2)
[1, 3, 5, 7, 9]
sage: range(10, 0, -1)
[10, 9, 8, 7, 6, 5, 4, 3, 2, 1]
sage: for i in range(4):
         print i, i*i
. . . . :
0 0
1 1
2 4
3 9
```

You can use the *continue* keyword to immediately go to the next item in the loop:

If you want to break out of the loop, use the *break* keyword:

If you need to keep track of both the position in the list and its value, one (not so elegant) way would be to do the following:

```
sage: l = ['a', 'b', 'c']
sage: for i in range(len(l)):
```

```
....: print i, l[i]
0 a
1 b
2 c
```

It's cleaner to use enumerate () which provides the index as well as the value:

```
sage: l = ['a', 'b', 'c']
sage: for i, letter in enumerate(l):
....: print i, letter
0 a
1 b
2 c
```

You could get a similar result to the result of the enumerate() function by using zip() to zip two lists together:

```
sage: l = ['a', 'b', 'c']
sage: for i, letter in zip(range(len(l)), l):
....: print i, letter
0 a
1 b
2 c
```

For loops work using Python's iterator protocol. This allows all sorts of different objects to be looped over. For example:

```
sage: for i in GF(5):
....: print i, i*i
0 0
1 1
2 4
3 4
4 1
```

How does this work?

```
sage: it = iter(GF(5)); it
<generator object __iter__ at 0x...>
sage: next(it)
0

sage: next(it)
1

sage: next(it)
2

sage: next(it)
3

sage: next(it)
4

sage: next(it)
4

sage: next(it)
5

sage: next(it)
4
Sage: next(it)

Traceback (most recent call last):
...
StopIteration
```

```
sage: R = GF(5)
sage: R.__iter___??
```

The command *yield* provides a very convenient way to produce iterators. We'll see more about it in a bit.

Exercises For each of the following sets, compute the list of its elements and their sum. Use two different ways, if possible: with a loop, and using a list comprehension.

1. The first n terms of the harmonic series:

$$\sum_{i=1}^{n} \frac{1}{i}$$

```
sage: # edit here
```

2. The odd integers between 1 and n:

```
sage: # edit here
```

3. The first n odd integers:

```
sage: # edit here
```

4. The integers between 1 and n that are neither divisible by 2 nor by 3 nor by 5:

```
sage: # edit here
```

5. The first n integers between 1 and n that are neither divisible by 2 nor by 3 nor by 5:

```
sage: # edit here
```

Functions

Functions are defined using the def statement, and values are returned using the return keyword:

```
sage: def f(x):
....: return x*x
sage: f(2)
```

Functions can be recursive:

```
sage: def fib(n):
....:     if n <= 1:
....:         return 1
....:     else:
....:         return fib(n-1) + fib(n-2)

sage: [fib(i) for i in range(10)]
[1, 1, 2, 3, 5, 8, 13, 21, 34, 55]</pre>
```

Functions are first class objects like any other. For example, they can be passed in as arguments to other functions:

```
sage: f
<function f at 0x...>
sage: def compose(f, x, n): # computes f(f(...f(x)))
....: for i in range(n):
x = f(x)
....: return x
                             # this change is local to this function call!
sage: compose(f, 2, 3)
256
sage: def add_one(x):
\dots: return x + 1
sage: compose(add_one, 2, 3)
You can give default values for arguments in functions:
sage: def add_n(x, n=1):
\dots: return x + n
sage: add_n(4)
sage: add_n(4, n=100)
104
sage: add_n(4, 1000)
1004
You can return multiple values from a function:
sage: def g(x):
....: return x, x*x
sage: g(2)
(2, 4)
sage: type(g)
<type 'function'>
sage: a,b = g(100)
sage: a
100
sage: b
10000
```

You can also take a variable number of arguments and keyword arguments in a function:

```
sage: def h(*args, **kwds):
....: print type(args), args
....: print type(kwds), kwds
```

```
sage: h(1,2,3,n=4)
<type 'tuple'> (1, 2, 3)
<type 'dict'> {'n': 4}
```

Let's use the *yield* instruction to make a generator for the Fibonacci numbers up to n:

```
sage: def fib_gen(n):
         if n < 1:
             return
         a = b = 1
. . . . :
         yield b
. . . . :
         while b < n:
             yield b
. . . . :
              a, b = b, b+a
sage: for i in fib_gen(50):
....: print i
1
2
3
5
8
13
21
34
```

Exercises

- 1. Write a function is even which returns True if n is even and False otherwise.
- 2. Write a function every_other which takes a list 1 as input and returns a list containing every other element of 1.
- 3. Write a generator every_other which takes an iterable 1 as input, and returns every other element of 1, one after the other.
- 4. Write a function which computes the n-th Fibonacci number. Try to improve performance.

Todo

- Definition of hashable
- Introduction to the debugger.

12.1.12 Tutorial: Comprehensions, Iterators, and Iterables

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List comprehensions

List comprehensions are a very handy way to construct lists in Python. You can use either of the following idioms:

```
[ <expr> for <name> in <iterable> ]
[ <expr> for <name> in <iterable> if <condition> ]
```

For example, here are some lists of squares:

```
sage: [ i^2 for i in [1, 3, 7] ]
[1, 9, 49]
sage: [ i^2 for i in range(1,10) ]
[1, 4, 9, 16, 25, 36, 49, 64, 81]
sage: [ i^2 for i in range(1,10) if i % 2 == 1]
[1, 9, 25, 49, 81]
```

And a variant on the latter:

```
sage: [i^2 if i ^2 2 == 1 else 2 for i in range(10)] [2, 1, 2, 9, 2, 25, 2, 49, 2, 81]
```

Exercises

1. Construct the list of the squares of the prime integers between 1 and 10:

```
sage: # edit here
```

2. Construct the list of the perfect squares less than 100 (hint: use srange() to get a list of Sage integers together with the method i.sqrtrem()):

```
sage: # edit here
```

One can use more than one iterable in a list comprehension:

```
sage: [ (i,j) for i in range(1,6) for j in range(1,i) ]
[(2, 1), (3, 1), (3, 2), (4, 1), (4, 2), (4, 3), (5, 1), (5, 2), (5, 3), (5, 4)]
```

Warning: Mind the order of the nested loop in the previous expression.

If instead one wants to build a list of lists, one can use nested lists as in:

```
sage: [ [ binomial(n, i) for i in range(n+1) ] for n in range(10) ]
[[1],
[1, 1],
[1, 2, 1],
[1, 3, 3, 1],
[1, 4, 6, 4, 1],
[1, 5, 10, 10, 5, 1],
[1, 6, 15, 20, 15, 6, 1],
[1, 7, 21, 35, 35, 21, 7, 1],
[1, 8, 28, 56, 70, 56, 28, 8, 1],
[1, 9, 36, 84, 126, 126, 84, 36, 9, 1]]
```

Exercises

1. Compute the list of pairs (i, j) of non negative integers such that i is at most 5, j is at most 8, and i and j are co-prime:

```
sage: # edit here
```

2. Compute the same list for i < j < 10:

```
sage: # edit here
```

Iterators

Definition

To build a comprehension, Python actually uses an *iterator*. This is a device which runs through a bunch of objects, returning one at each call to the next method. Iterators are built using parentheses:

```
sage: it = (binomial(8, i) for i in range(9))
sage: next(it)
1

sage: next(it)
8
sage: next(it)
28
sage: next(it)
```

You can get the list of the results that are not yet *consumed*:

```
sage: list(it)
[70, 56, 28, 8, 1]
```

Asking for more elements triggers a StopIteration exception:

```
sage: next(it)
Traceback (most recent call last):
...
StopIteration
```

An iterator can be used as argument for a function. The two following idioms give the same results; however, the second idiom is much more memory efficient (for large examples) as it does not expand any list in memory:

```
sage: sum( [ binomial(8, i) for i in range(9) ] )
256
sage: sum( binomial(8, i) for i in xrange(9) )
256
```

Exercises

```
1. Compute the sum of \binom{10}{i} for all even i:
```

```
sage: # edit here
```

2. Compute the sum of the gcd's of all co-prime numbers i,j for i < j < 10:

```
sage: # edit here
```

Typical usage of iterators

Iterators are very handy with the functions all(), any(), and exists():

```
sage: all([True, True, True, True])
True
```

```
sage: all([True, False, True, True])
False
sage: any([False, False, False])
sage: any([False, False, True, False])
True
Let's check that all the prime numbers larger than 2 are odd:
sage: all( is_odd(p) for p in range(1,100) if is_prime(p) and p>2 )
True
It is well know that if 2^p-1 is prime then p is prime:
sage: def mersenne(p): return 2^p -1
sage: [ is_prime(p) for p in range(20) if is_prime(mersenne(p)) ]
[True, True, True, True, True, True]
The converse is not true:
sage: all( is_prime(mersenne(p)) for p in range(1000) if is_prime(p) )
False
Using a list would be much slower here:
sage: %time all( is_prime(mersenne(p)) for p in range(1000) if is_prime(p) ) # not tested
CPU times: user 0.00 s, sys: 0.00 s, total: 0.00 s
Wall time: 0.00 s
False
sage: %time all( [ is_prime(mersenne(p)) for p in range(1000) if is_prime(p)] ) # not tested
CPU times: user 0.72 s, sys: 0.00 s, total: 0.73 s
Wall time: 0.73 s
False
You can get the counterexample using exists (). It takes two arguments: an iterator and a function which tests the
property that should hold:
sage: exists( (p for p in range(1000) if is_prime(p)), lambda p: not is_prime(mersenne(p)) )
(True, 11)
An alternative way to achieve this is:
sage: counter_examples = (p for p in range(1000) if is_prime(p) and not is_prime(mersenne(p)))
sage: next(counter_examples)
 Exercises
   1. Build the list \{i^3 \mid -10 < i < 10\}. Can you find two of those cubes u and v such that u + v = 218?
      sage: # edit here
```

itertools

At its name suggests itertools is a module which defines several handy tools for manipulating iterators:

```
sage: 1 = [3, 234, 12, 53, 23]
sage: [(i, 1[i]) for i in range(len(1))]
[(0, 3), (1, 234), (2, 12), (3, 53), (4, 23)]
```

The same results can be obtained using enumerate():

```
sage: list(enumerate(1))
[(0, 3), (1, 234), (2, 12), (3, 53), (4, 23)]
```

Here is the analogue of list slicing:

```
sage: list(Permutations(3))
[[1, 2, 3], [1, 3, 2], [2, 1, 3], [2, 3, 1], [3, 1, 2], [3, 2, 1]]
sage: list(Permutations(3))[1:4]
[[1, 3, 2], [2, 1, 3], [2, 3, 1]]
sage: import itertools
sage: list(itertools.islice(Permutations(3), 1, 4))
[[1, 3, 2], [2, 1, 3], [2, 3, 1]]
```

The functions map() and filter() also have an analogue:

```
sage: list(itertools.imap(lambda z: z.cycle_type(), Permutations(3)))
[[1, 1, 1], [2, 1], [2, 1], [3], [3], [2, 1]]

sage: list(itertools.ifilter(lambda z: z.has_pattern([1,2]), Permutations(3)))
[[1, 2, 3], [1, 3, 2], [2, 1, 3], [2, 3, 1], [3, 1, 2]]
```

Exercises

1. Define an iterator for the *i*-th prime for 5 < i < 10:

```
sage: # edit here
```

Defining new iterators

One can very easily write new iterators using the keyword yield. The following function does nothing interesting beyond demonstrating the use of yield:

```
sage: def f(n):
....: for i in range(n):
....: yield i
sage: [ u for u in f(5) ]
[0, 1, 2, 3, 4]
```

Iterators can be recursive:

```
sage: def words(alphabet,1):
....:    if 1 == 0:
....:        yield[]
....:    else:
```

```
for word in words(alphabet, 1-1):
    for a in alphabet:
        yield word + [a]

sage: [ w for w in words(['a','b','c'], 3) ]
[['a', 'a', 'a'], ['a', 'a', 'b'], ['a', 'a', 'c'], ['a', 'b', 'a'], ['a', 'b', 'b'], ['a', 'b', 'c']
sage: sum(1 for w in words(['a','b','c'], 3))
27
```

Here is another recursive iterator:

```
sage: def dyck_words(1):
\dots: if l==0:
              yield ''
. . . . :
. . . . :
         else:
...:
              for k in range(l):
                   for w1 in dyck_words(k):
                       for w2 in dyck_words(1-k-1):
. . . . :
                           yield '('+w1+')'+w2
. . . . :
sage: list(dyck_words(4))
['()()()()',
′()()(())′,
′()(())()′,
′()(()())′,
′()(((()))′,
′(())()()′,
′(())(())′,
′(()())()′,
′(((()))()′,
′(()()())′,
′(()(()))′,
′(((())())′,
'(((()()))',
′((((())))']
sage: sum(1 for w in dyck_words(5))
42
```

Exercises

1. Write an iterator with two parameters n, l iterating through the set of nondecreasing lists of integers smaller than n of length l:

```
sage: # edit here
```

Standard Iterables

Finally, many standard Python and Sage objects are iterable; that is one may iterate through their elements:

```
sage: sum( x^{1}en(s) for s in Subsets(8) ) x^{8} + 8*x^{7} + 28*x^{6} + 56*x^{5} + 70*x^{4} + 56*x^{3} + 28*x^{2} + 8*x + 1 sage: sum( x^{1}ength() for p in Permutations(3) ) x^{3} + 2*x^{2} + 2*x + 1
```

```
sage: factor(sum( x^p.length() for p in Permutations(3) ))
(x^2 + x + 1) * (x + 1)
sage: P = Permutations(5)
sage: all( p in P for p in P )
True
sage: for p in GL(2, 2): print p; print
[1 0]
[0 1]
[0 1]
[1 0]
[0 1]
[1 1]
[1 1]
[0 1]
[1 1]
[1 0]
[1 0]
[1 1]
sage: for p in Partitions(3): print p
[3]
[2, 1]
[1, 1, 1]
Beware of infinite loops:
sage: for p in Partitions(): print p
                                                 # not tested
sage: for p in Primes(): print p
                                                  # not tested
Infinite loops can nevertheless be very useful:
sage: exists( Primes(), lambda p: not is_prime(mersenne(p)) )
(True, 11)
sage: counter_examples = (p for p in Primes() if not is_prime(mersenne(p)))
sage: next(counter_examples)
11
```

12.1.13 Tutorial: Objects and Classes in Python and Sage

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This tutorial is an introduction to object-oriented programming in Python and Sage. It requires basic knowledge about imperative/procedural programming (the most common programming style) – that is, conditional instructions, loops, functions (see the "Programming" section of the Sage tutorial) – but no further knowledge about objects and classes

is assumed. It is designed as an alternating sequence of formal introduction and exercises. *Solutions to the exercises* are given at the end.

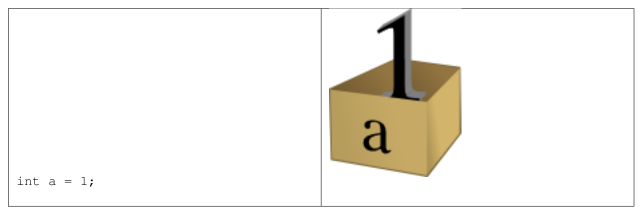
Foreword: variables, names and objects

As an object-oriented language, Python's 'variables' behavior may be surprising for people used to imperative languages like C or Maple. The reason is that they are **not variables but names**.

The following explanation is borrowed from David Goodger.

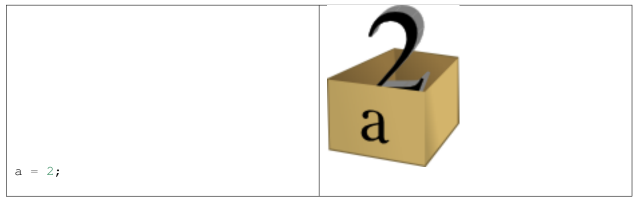
Other languages have "variables"

In many other languages, assigning to a variable puts a value into a box.



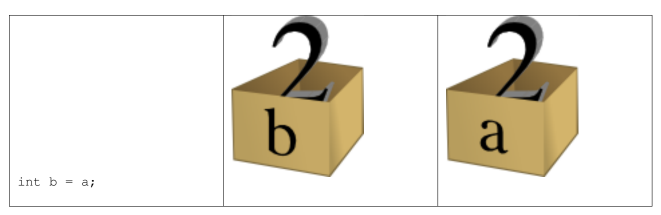
Box "a" now contains an integer 1.

Assigning another value to the same variable replaces the contents of the box:



Now box "a" contains an integer 2.

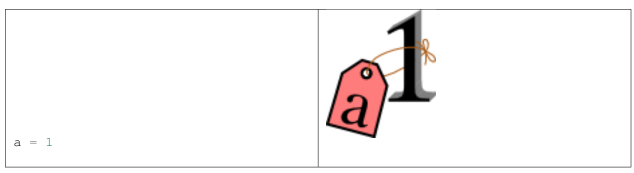
Assigning one variable to another makes a copy of the value and puts it in the new box:



"b" is a second box, with a copy of integer 2. Box "a" has a separate copy.

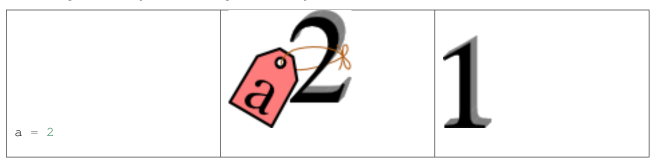
Python has "names"

In Python, a "name" or "identifier" is like a parcel tag (or nametag) attached to an object.



Here, an integer 1 object has a tag labelled "a".

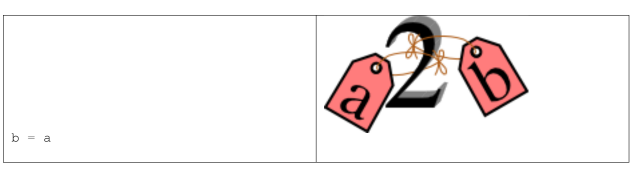
If we reassign to "a", we just move the tag to another object:



Now the name "a" is attached to an integer 2 object.

The original integer 1 object no longer has a tag "a". It may live on, but we can't get to it through the name "a". (When an object has no more references or tags, it is removed from memory.)

If we assign one name to another, we're just attaching another nametag to an existing object:



The name "b" is just a second tag bound to the same object as "a".

Although we commonly refer to "variables" even in Python (because it's common terminology), we really mean "names" or "identifiers". In Python, "variables" are nametags for values, not labelled boxes.

Warning: As a consequence, when there are **two tags** "a" and "b" on the **same object**, modifying the object tagged "b" also modifies the object tagged "a":

```
sage: a = [1,2,3]
sage: b = a
sage: b[1] = 0
sage: a
[1, 0, 3]
```

Note that reassigning the tag "b" (rather than modifying the object with that tag) doesn't affect the object tagged "a":

```
sage: b = 7
sage: b
7
sage: a
[1, 0, 3]
```

Object-oriented programming paradigm

The **object-oriented programming** paradigm relies on the two following fundamental rules:

- 1. Anything of the real (or mathematical) world which needs to be manipulated by the computer is modeled by an **object**.
- 2. Each object is an instance of some class.

At this point, those two rules are a little meaningless, so let's give some more or less precise definitions of the terms:

object a **portion of memory** which contains the information needed to model the real world thing.

class defines the data structure used to store the objects which are instances of the class together with their behavior.

Let's start with some examples: We consider the vector space over **Q** whose basis is indexed by permutations, and a particular element in it:

```
sage: F = CombinatorialFreeModule(QQ, Permutations())
sage: el = 3*F([1,3,2])+ F([1,2,3])
sage: el
B[[1, 2, 3]] + 3*B[[1, 3, 2]]
```

(For each permutation, say [1, 3, 2], the corresponding element in F is denoted by B[[1, 3, 2]] - in a CombinatorialFreeModule, if an element is indexed by x, then by default its print representation is B[x].)

In Python, everything is an object so there isn't any difference between types and classes. One can get the class of the object el by:

```
sage: type(el)
<class 'sage.combinat.free_module.CombinatorialFreeModule_with_category.element_class'>
```

As such, this is not very informative. We'll come back to it later. The data associated to objects are stored in so-called **attributes**. They are accessed through the syntax obj.attribute_name. For an element of a combinatorial free module, the main attribute is called _monomial_coefficients. It is a dictionary associating coefficients to indices:

```
sage: el._monomial_coefficients
{[1, 2, 3]: 1, [1, 3, 2]: 3}
```

Modifying the attribute modifies the objects:

```
sage: el._monomial_coefficients[Permutation([3,2,1])] = 1/2
sage: el
B[[1, 2, 3]] + 3*B[[1, 3, 2]] + 1/2*B[[3, 2, 1]]
```

Warning: as a user, you are *not* supposed to do such a modification by yourself (see note on *private attributes* below).

As an element of a vector space, el has a particular behavior:

```
sage: 2*el
2*B[[1, 2, 3]] + 6*B[[1, 3, 2]] + B[[3, 2, 1]]
sage: sorted(el.support())
[[1, 2, 3], [1, 3, 2], [3, 2, 1]]
sage: el.coefficient([1, 2, 3])
1
```

The behavior is defined through **methods** (support, coefficient). Note that this is true even for equality, printing or mathematical operations. For example, the call a == b actually is translated to the method call a.__eq__(b). The names of those special methods which are usually called through operators are fixed by the Python language and are of the form __name__. Examples include __eq__ and __le__ for operators == and <=, __repr__ (see <u>Sage specifics about classes</u>) for printing, __add__ and __mult__ for operators + and *. See http://docs.python.org/library/ for a complete list.

```
sage: el.__eq__(F([1,3,2]))
False
sage: el.__repr__()
'B[[1, 2, 3]] + 3*B[[1, 3, 2]] + 1/2*B[[3, 2, 1]]'
sage: el.__mul__(2)
2*B[[1, 2, 3]] + 6*B[[1, 3, 2]] + B[[3, 2, 1]]
```

Some particular actions modify the data structure of el:

```
sage: el.rename("bla")
sage: el
bla
```

Note: The class is stored in a particular attribute called __class__, and the normal attributes are stored in a dictionary called __dict__:

Lots of Sage objects are not Python objects but compiled Cython objects. Python sees them as builtin objects and you don't have access to the data structure. Examples include integers and permutation group elements:

Note: Each object corresponds to a portion of memory called its **identity** in Python. You can get the identity using id:

```
sage: el = Integer(9)
sage: id(el) # random
139813642977744
sage: el1 = el; id(el1) == id(el)
True
sage: el1 is el
True
```

In Python (and therefore in Sage), two objects with the same identity will be equal, but the converse is not true in general. Thus the identity function is different from mathematical identity:

```
sage: e12 = Integer(9)
sage: e12 == e11
True
sage: e12 is e11
False
sage: id(e12) == id(e1)
False
```

Summary

To define some object, you first have to write a **class**. The class will define the methods and the attributes of the object. **method** particular kind of function associated with an object used to get information about the object or to manipulate

attribute variable where information about the object is stored.

An example: glass of beverage in a restaurant

Let's write a small class about glasses in a restaurant:

```
sage: class Glass(object):
          def __init__(self, size):
. . . . :
               assert size > 0
. . . . :
               self._size = float(size) # an attribute
. . . . :
               self._content = float(0.0) # another attribute
. . . . :
                __repr__(self):
. . . . :
               if self._content == 0.0:
                   return "An empty glass of size %s"%(self._size)
               else:
. . . . :
                   return "A glass of size %s cl containing %s cl of water"%(
. . . . :
                            self._size, self._content)
. . . . :
          def fill(self):
. . . . :
               self._content = self._size
          def empty(self):
               self.\_content = float(0.0)
```

Let's create a small glass:

```
sage: myGlass = Glass(10); myGlass
An empty glass of size 10.0
sage: myGlass.fill(); myGlass
A glass of size 10.0 cl containing 10.0 cl of water
sage: myGlass.empty(); myGlass
An empty glass of size 10.0
```

Some comments:

- 1. The definition of the class Glass defines two attributes, _size and _content. It defines four methods, __init__, __repr__, fill, and empty. (Any instance of this class will also have other attributes and methods, inherited from the class object. See *Inheritance* below.)
- 2. The method __init__ is used to initialize the object: it is used by the so-called **constructor** of the class that is executed when calling Glass (10).
- 3. The method __repr__ returns a string which is used to print the object, for example in this case when evaluating myGlass.

Note: Private Attributes

• Most of the time, in order to ensure consistency of the data structures, the user is not supposed to directly change certain attributes of an object. Those attributes are called **private**. Since there is no mechanism to ensure privacy in Python, the convention is the following: private attributes have names beginning with an underscore.

- As a consequence, attribute access is only made through methods. Methods for reading or writing a private attribute are called accessors.
- Methods which are only for internal use are also prefixed with an underscore.

Exercises

- 1. Add a method is_empty which returns true if a glass is empty.
- 2. Define a method drink with a parameter amount which allows one to partially drink the water in the glass. Raise an error if one asks to drink more water than there is in the glass or a negative amount of water.
- 3. Allows the glass to be filled with wine, beer or another beverage. The method fill should accept a parameter beverage. The beverage is stored in an attribute _beverage. Update the method __repr__ accordingly.
- 4. Add an attribute _clean and methods is_clean and wash. At the creation a glass is clean, as soon as it's filled it becomes dirty, and it must be washed to become clean again.
- 5. Test everything.
- 6. Make sure that everything is tested.
- 7. Test everything again.

Inheritance

The problem: objects of **different** classes may share a **common behavior**.

For example, if one wants to deal with different dishes (forks, spoons, ...), then there is common behavior (becoming dirty and being washed). So the different classes associated to the different kinds of dishes should have the same clean, is_clean and wash methods. But copying and pasting code is very bad for maintenance: mistakes are copied, and to change anything one has to remember the location of all the copies. So there is a need for a mechanism which allows the programmer to factorize the common behavior. It is called **inheritance** or **sub-classing**: one writes a base class which factorizes the common behavior and then reuses the methods from this class.

We first write a small class "AbstractDish" which implements the "clean-dirty-wash" behavior:

```
sage: class AbstractDish(object):
          def __init__(self):
. . . . :
               self._clean = True
          def is_clean(self):
. . . . :
               return self._clean
. . . . :
          def state(self):
. . . . :
               return "clean" if self.is_clean() else "dirty"
. . . . :
          def __repr__(self):
               return "An unspecified %s dish"%self.state()
          def _make_dirty(self):
               self._clean = False
          def wash(self):
. . . . :
               self._clean = True
```

Now one can reuse this behavior within a class Spoon:

```
sage: class Spoon(AbstractDish): # Spoon inherits from AbstractDish
...: def __repr__(self):
...: return "A %s spoon"%self.state()
...: def eat_with(self):
...: self._make_dirty()
```

Let's test it:

```
sage: s = Spoon(); s
A clean spoon
sage: s.is_clean()
True
sage: s.eat_with(); s
A dirty spoon
sage: s.is_clean()
False
sage: s.wash(); s
A clean spoon
```

Summary

- Any class can reuse the behavior of another class. One says that the subclass inherits from the superclass or that it derives from it.
- 2. Any instance of the subclass is also an instance of its superclass:

```
sage: type(s)
<class '__main__.Spoon'>
sage: isinstance(s, Spoon)
True
sage: isinstance(s, AbstractDish)
True
```

3. If a subclass redefines a method, then it replaces the former one. One says that the subclass **overloads** the method. One can nevertheless explicitly call the hidden superclass method.

```
sage: s.__repr__()
'A clean spoon'
sage: Spoon.__repr__(s)
'A clean spoon'
sage: AbstractDish.__repr__(s)
'An unspecified clean dish'
```

Note: Advanced superclass method call

Sometimes one wants to call an overloaded method without knowing in which class it is defined. To do this, use the super operator:

```
sage: super(Spoon, s).__repr__()
'An unspecified clean dish'
```

A very common usage of this construct is to call the __init__ method of the superclass:

```
sage: class Spoon(AbstractDish):
....:     def __init__(self):
....:         print "Building a spoon"
....:         super(Spoon, self).__init__()
....:     def __repr__(self):
....:         return "A %s spoon"%self.state()
....:         def eat_with(self):
....:         self._make_dirty()
sage: s = Spoon()
Building a spoon
```

```
sage: s
A clean spoon
```

Exercises

- 1. Modify the class Glasses so that it inherits from Dish.
- 2. Write a class Plate whose instance can contain any meal together with a class Fork. Avoid as much as possible code duplication (hint: you can write a factorized class ContainerDish).
- 3. Test everything.

Sage specifics about classes

Compared to Python, Sage has particular ways to handle objects:

- Any classes for mathematical objects in Sage should inherit from SageObject rather than from object. Most of the time, they actually inherit from a subclass such as Parent or Element.
- Printing should be done through _repr_ instead of __repr_ to allow for renaming.
- More generally, Sage-specific special methods are usually named _meth_ rather than __meth__. For example, lots of classes implement _hash_ which is used and cached by __hash__. In the same vein, elements of a group usually implement _mul_, so that there is no need to take care about coercions as they are done in mul_.

For more details, see the Sage Developer's Guide.

Solutions to the exercises

1. Here is a solution to the first exercise:

```
sage: class Glass(object):
          def __init__(self, size):
. . . . :
               assert size > 0
. . . . :
               self._size = float(size)
. . . . :
               self.wash()
          def __repr__(self):
               if self._content == 0.0:
                    return "An empty glass of size %s"% (self._size)
. . . . :
               else:
. . . . :
                   return "A glass of size %s cl containing %s cl of %s"%(
. . . . :
                             self._size, self._content, self._beverage)
. . . . :
          def content (self):
               return self._content
          def beverage(self):
. . . . :
               return self._beverage
. . . . :
          def fill(self, beverage = "water"):
. . . . :
               if not self.is_clean():
. . . . :
                   raise ValueError("Don't want to fill a dirty glass")
               self._clean = False
               self._content = self._size
               self._beverage = beverage
. . . . :
          def empty(self):
. . . . :
               self._content = float(0.0)
. . . . :
```

```
def is_empty(self):
. . . . :
              return self._content == 0.0
. . . . :
          def drink(self, amount):
. . . . :
               if amount <= 0.0:
. . . . :
. . . . :
                    raise ValueError("amount must be positive")
               elif amount > self._content:
. . . . :
                   raise ValueError("not enough beverage in the glass")
. . . . :
               else:
. . . . :
                   self._content -= float(amount)
. . . . :
         def is_clean(self):
. . . . :
              return self._clean
. . . . :
          def wash(self):
. . . . :
               self.\_content = float(0.0)
. . . . :
               self._beverage = None
. . . . :
               self._clean = True
. . . . :
```

2. Let's check that everything is working as expected:

```
sage: G = Glass(10.0)
sage: G
An empty glass of size 10.0
sage: G.is_empty()
True
sage: G.drink(2)
Traceback (most recent call last):
ValueError: not enough beverage in the glass
sage: G.fill("beer")
sage: G
A glass of size 10.0 cl containing 10.0 cl of beer
sage: G.is_empty()
False
sage: G.is_clean()
False
sage: G.drink(5.0)
A glass of size 10.0 cl containing 5.0 cl of beer
sage: G.is_empty()
False
sage: G.is_clean()
False
sage: G.drink(5)
sage: G
An empty glass of size 10.0
sage: G.is_clean()
False
sage: G.fill("orange juice")
Traceback (most recent call last):
ValueError: Don't want to fill a dirty glass
sage: G.wash()
sage: G
An empty glass of size 10.0
sage: G.fill("orange juice")
sage: G
A glass of size 10.0 cl containing 10.0 cl of orange juice
```

3. Here is the solution to the second exercice:

```
sage: class AbstractDish(object):
         def __init__(self):
. . . . :
               self._clean = True
. . . . :
          def is_clean(self):
. . . . :
. . . . :
               return self._clean
         def state(self):
. . . . :
              return "clean" if self.is_clean() else "dirty"
. . . . :
         def __repr__(self):
. . . . :
              return "An unspecified %s dish"%self.state()
. . . . :
         def _make_dirty(self):
. . . . :
              self._clean = False
. . . . :
         def wash(self):
. . . . :
               self._clean = True
. . . . :
sage: class ContainerDish(AbstractDish):
         def __init__(self, size):
. . . . :
. . . . :
               assert size > 0
               self._size = float(size)
. . . . :
               self._content = float(0)
. . . . :
               super(ContainerDish, self).__init__()
. . . . :
         def content(self):
. . . . :
              return self._content
. . . . :
         def empty(self):
. . . . :
              self.\_content = float(0.0)
. . . . :
         def is_empty(self):
. . . . :
              return self._content == 0.0
. . . . :
         def wash(self):
. . . . :
               self._content = float(0.0)
. . . . :
               super(ContainerDish, self).wash()
. . . . :
sage: class Glass(ContainerDish):
....: def __repr__(self):
              if self._content == 0.0:
. . . . :
                   return "An empty glass of size %s"% (self._size)
. . . . :
               else:
. . . . :
                    return "A glass of size %s cl containing %s cl of %s"%(
. . . . :
. . . . :
                             self._size, self._content, self._beverage)
         def beverage(self):
. . . . :
               return self._beverage
. . . . :
         def fill(self, beverage = "water"):
. . . . :
               if not self.is_clean():
. . . . :
                    raise ValueError("Don't want to fill a dirty glass")
. . . . :
               self._make_dirty()
               self._content = self._size
. . . . :
               self._beverage = beverage
. . . . :
         def drink(self, amount):
. . . . :
              if amount <= 0.0:
. . . . :
                   raise ValueError("amount must be positive")
. . . . :
. . . . :
               elif amount > self._content:
                   raise ValueError("not enough beverage in the glass")
. . . . :
               else:
. . . . :
                   self._content -= float(amount)
. . . . :
         def wash(self):
. . . . :
               self._beverage = None
. . . . :
               super(Glass, self).wash()
```

4. Let's check that everything is working as expected:

```
sage: G = Glass(10.0)
sage: G
An empty glass of size 10.0
sage: G.is_empty()
True
sage: G.drink(2)
Traceback (most recent call last):
ValueError: not enough beverage in the glass
sage: G.fill("beer")
sage: G
A glass of size 10.0 cl containing 10.0 cl of beer
sage: G.is_empty()
False
sage: G.is_clean()
False
sage: G.drink(5.0)
sage: G
A glass of size 10.0 cl containing 5.0 cl of beer
sage: G.is_empty()
False
sage: G.is_clean()
False
sage: G.drink(5)
sage: G
An empty glass of size 10.0
sage: G.is_clean()
sage: G.fill("orange juice")
Traceback (most recent call last):
ValueError: Don't want to fill a dirty glass
sage: G.wash()
sage: G
An empty glass of size 10.0
sage: G.fill("orange juice")
sage: G
A glass of size 10.0 cl containing 10.0 cl of orange juice
```

Todo

give the example of the class Plate.

That all folks!

12.1.14 Functional Programming for Mathematicians

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This tutorial discusses some techniques of functional programming that might be of interest to mathematicians or people who use Python for scientific computation. We start off with a brief overview of procedural and object-oriented programming, and then discuss functional programming techniques. Along the way, we briefly review Python's built-in support for functional programming, including filter, lambda, map and reduce. The tutorial concludes with some resources on detailed information on functional programming using Python.

Styles of programming

Python supports several styles of programming. You could program in the procedural style by writing a program as a list of instructions. Say you want to implement addition and multiplication over the integers. A procedural program to do so would be as follows:

The Python module operator defines several common arithmetic and comparison operators as named functions. Addition is defined in the built-in function operator.add and multiplication is defined in operator.mul. The above example can be worked through as follows:

```
sage: from operator import add
sage: from operator import mul
sage: add(2, 3)
5
sage: mul(2, 3)
6
```

Another common style of programming is called object-oriented programming. Think of an object as code that encapsulates both data and functionalities. You could encapsulate integer addition and multiplication as in the following object-oriented implementation:

```
sage: class MyInteger:
....:     def __init__(self):
....:         self.cardinality = "infinite"
....:     def add(self, a, b):
....:         return a + b
....:     def mult(self, a, b):
....:         return a * b
...
sage: myZZ = MyInteger()
sage: myZZ.cardinality
'infinite'
sage: myZZ.add(2, 3)
5
sage: myZZ.mult(2, 3)
```

Functional programming using map

Functional programming is yet another style of programming in which a program is decomposed into various functions. The Python built-in functions map, reduce and filter allow you to program in the functional style. The function

```
map(func, seq1, seq2, ...)
```

takes a function func and one or more sequences, and apply func to elements of those sequences. In particular, you end up with a list like so:

```
[func(seq1[0], seq2[0], ...), func(seq1[1], seq2[1], ...), ...]
```

In many cases, using map allows you to express the logic of your program in a concise manner without using list comprehension. For example, say you have two lists of integers and you want to add them element-wise. A list comprehension to accomplish this would be as follows:

```
sage: A = [1, 2, 3, 4]
sage: B = [2, 3, 5, 7]
sage: [A[i] + B[i] for i in range(len(A))]
[3, 5, 8, 11]
```

Alternatively, you could use the Python built-in addition function operator. add together with map to achieve the same result:

```
sage: from operator import add
sage: A = [1, 2, 3, 4]
sage: B = [2, 3, 5, 7]
sage: map(add, A, B)
[3, 5, 8, 11]
```

An advantage of map is that you do not need to explicitly define a for loop as was done in the above list comprehension.

Define small functions using lambda

There are times when you want to write a short, one-liner function. You could re-write the above addition function as follows:

```
sage: def add_ZZ(a, b): return a + b
...
```

Or you could use a lambda statement to do the same thing in a much clearer style. The above addition and multiplication functions could be written using lambda as follows:

```
sage: add_ZZ = lambda a, b: a + b
sage: mult_ZZ = lambda a, b: a * b
sage: add_ZZ(2, 3)
5
sage: mult_ZZ(2, 3)
```

Things get more interesting once you combine map with the lambda statement. As an exercise, you might try to write a simple function that implements a constructive algorithm for the Chinese Remainder Theorem. You could use list comprehension together with map and lambda as shown below. Here, the parameter A is a list of integers and M is a list of moduli.

```
sage: x = crt(A, M); x
11
sage: mod(x, 3)
2
sage: mod(x, 4)
3
sage: mod(x, 5)
1
```

To produce a random matrix over a ring, say \mathbf{Z} , you could start by defining a matrix space and then obtain a random element of that matrix space:

```
sage: MS = MatrixSpace(ZZ, nrows=5, ncols=3)
sage: MS.random_element() # random

[ 6  1  0]
[-1  5  0]
[-1  0  0]
[-5  0  1]
[ 1 -1 -3]
```

Or you could use the function random_matrix:

```
sage: random_matrix(ZZ, nrows=5, ncols=3) # random
[ 2 -50   0]
[ -1   0   -6]
[ -4   -1   -1]
[  1   1   3]
[  2   -1   -1]
```

The next example uses map to construct a list of random integer matrices:

```
sage: rows = [randint(1, 10) for i in range(10)]
sage: cols = [randint(1, 10) for i in range(10)]
sage: rings = [ZZ]*10
sage: M = map(random_matrix, rings, rows, cols)
sage: M[0] # random
[ -1 -3 -1 -37 1 -1 -4 5]
[ 2 1 1 5 2 1 -2 1]
[ -1 0 -4 0 -2 1 -2 1]
```

If you want more control over the entries of your matrices than the random_matrix function permits, you could use lambda together with map as follows:

```
sage: rand_row = lambda n: [randint(1, 10) for i in range(n)]
sage: rand_mat = lambda nrows, ncols: [rand_row(ncols) for i in range(nrows)]
sage: matrix(rand_mat(5, 3)) # random

[ 2  9 10]
[ 8  8  9]
[ 6  7  6]
[ 9  2 10]
[ 2  6  2]
sage: rows = [randint(1, 10) for i in range(10)]
sage: cols = [randint(1, 10) for i in range(10)]
sage: M = map(rand_mat, rows, cols)
```

```
sage: M = map(matrix, M)
sage: M[0] # random
[ 9 1 5 2 10 10 1]
[ 3 4 3 7 4 3
[ 4 8
      7
        6 4
            2 101
     3 3 6 2
[ 1
   6
               1]
[5526434]
[6629451]
[10 2 5 5 7 10 4]
[2 7 3 5 10 8 1]
[1 5 1 7 8 8 6]
```

Reducing a sequence to a value

The function reduce takes a function of two arguments and apply it to a given sequence to reduce that sequence to a single value. The function sum is an example of a reduce function. The following sample code uses reduce and the built-in function operator.add to add together all integers in a given list. This is followed by using sum to accomplish the same task:

```
sage: from operator import add
sage: L = [1, 2, 3, 4, 5, 6, 7, 8, 9, 10]
sage: reduce(add, L)
55
sage: sum(L)
55
```

In the following sample code, we consider a vector as a list of real numbers. The dot product is then implemented using the functions operator.add and operator.mul, in conjunction with the built-in Python functions reduce and map. We then show how sum and map could be combined to produce the same result.

```
sage: from operator import add
sage: from operator import mul
sage: U = [1, 2, 3]
sage: V = [2, 3, 5]
sage: reduce(add, map(mul, U, V))
23
sage: sum(map(mul, U, V))
```

Or you could use Sage's built-in support for the dot product:

```
sage: u = vector([1, 2, 3])
sage: v = vector([2, 3, 5])
sage: u.dot_product(v)
23
```

Here is an implementation of the Chinese Remainder Theorem without using sum as was done previously. The version below uses operator.add and defines mul3 to multiply three numbers instead of two.

```
sage: def crt(A, M):
....:     from operator import add
....:     Mprod = prod(M)
....:     Mdiv = map(lambda x: Integer(Mprod / x), M)
....:     X = map(inverse_mod, Mdiv, M)
....:     mul3 = lambda a, b, c: a * b * c
```

Filtering with filter

The Python built-in function filter takes a function of one argument and a sequence. It then returns a list of all those items from the given sequence such that any item in the new list results in the given function returning True. In a sense, you are filtering out all items that satisfy some condition(s) defined in the given function. For example, you could use filter to filter out all primes between 1 and 50, inclusive.

```
sage: filter(is_prime, [1..50])
[2, 3, 5, 7, 11, 13, 17, 19, 23, 29, 31, 37, 41, 43, 47]
```

For a given positive integer n, the Euler phi function counts the number of integers a, with $1 \le a \le n$, such that gcd(a, n) = 1. You could use list comprehension to obtain all such a's when n = 20:

```
sage: [k for k in range(1, 21) if gcd(k, 20) == 1]
[1, 3, 7, 9, 11, 13, 17, 19]
```

A functional approach is to use lambda to define a function that determines whether or not a given integer is relatively prime to 20. Then you could use filter instead of list comprehension to obtain all the required a's.

```
sage: is_coprime = lambda k: gcd(k, 20) == 1
sage: filter(is_coprime, range(1, 21))
[1, 3, 7, 9, 11, 13, 17, 19]
```

The function primroots defined below returns all primitive roots modulo a given positive prime integer p. It uses filter to obtain a list of integers between 1 and p-1, inclusive, each integer in the list being relatively prime to the order of the multiplicative group $(\mathbf{Z}/p\mathbf{Z})^*$.

```
sage: def primroots(p):
....: g = primitive_root(p)
. . . . :
         znorder = p - 1
        is_coprime = lambda x: gcd(x, znorder) == 1
        good_odd_integers = filter(is_coprime, [1..p-1, step=2])
        all_primroots = [power_mod(g, k, p) for k in good_odd_integers]
        all_primroots.sort()
        return all_primroots
sage: primroots(3)
sage: primroots(5)
sage: primroots(7)
[3, 5]
sage: primroots(11)
[2, 6, 7, 8]
sage: primroots(13)
[2, 6, 7, 11]
sage: primroots(17)
[3, 5, 6, 7, 10, 11, 12, 14]
```

```
sage: primroots(23)
[5, 7, 10, 11, 14, 15, 17, 19, 20, 21]
sage: primroots(29)
[2, 3, 8, 10, 11, 14, 15, 18, 19, 21, 26, 27]
sage: primroots(31)
[3, 11, 12, 13, 17, 21, 22, 24]
```

Further resources

This has been a rather short tutorial to functional programming with Python. The Python standard documentation has a list of built-in functions, many of which are useful in functional programming. For example, you might want to read up on all, any, max, min, and zip. The Python module operator has numerous built-in arithmetic and comparison operators, each operator being implemented as a function whose name reflects its intended purpose. For arithmetic and comparison operations, it is recommended that you consult the operator module to determine if there is a built-in function that satisfies your requirement. The module itertools has numerous built-in functions to efficiently process sequences of items. The functions filter, map and zip have their counterparts in itertools as itertools.ifilter, itertools.imap and itertools.izip.

Another useful resource for functional programming in Python is the Functional Programming HOWTO by A. M. Kuchling. Steven F. Lott's book Building Skills in Python has a chapter on Functional Programming using Collections. See also the chapter Functional Programming from Mark Pilgrim's book Dive Into Python.

You might also want to consider experimenting with Haskell for expressing mathematical concepts. For an example of Haskell in expressing mathematical algorithms, see J. Gibbons' article Unbounded Spigot Algorithms for the Digits of Pi in the American Mathematical Monthly.

12.1.15 How to implement new algebraic structures in Sage

Contents

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Sage's category and coercion framework

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The aim of this tutorial is to explain how one can benefit from Sage's category framework and coercion model when implementing new algebraic structures. It is based on a worksheet created in 2011.

We illustrate the concepts of Sage's category framework and coercion model by means of a detailed example, namely a toy implementation of fraction fields. The code is developed step by step, so that the reader can focus on one detail in each part of this tutorial. The complete code can be found in the appendix.

Outline

Use existing base classes

For using Sage's coercion system, it is essential to work with sub-classes of sage.structure.parent.Parent or sage.structure.element.Element, respectively. They provide default implementations of many "magical" double-underscore Python methods, which must not be overridden. Instead, the actual implementation should be in *single underscore* methods, such as <code>_add_</code> or <code>_mul_</code>.

• Turn your parent structure into an object of a category

Declare the category during initialisation—Your parent structure will inherit further useful methods and consistency tests.

• Provide your parent structure with an element class

Assign to it an attribute called Element—The elements will inherit further useful methods from the category. In addition, some basic conversions will immediately work.

• Implement further conversions

Never override a parent's __call__ method! Provide the method_element_constructor_instead.

Declare coercions

If a conversion happens to be a morphism, you may consider to turn it into a coercion. It will then *implicitly* be used in arithmetic operations.

• Advanced coercion: Define construction functors for your parent structure

Sage will automatically create new parents for you when needed, by the so-called sage.categories.pushout.pushout() construction.

• Run the automatic test suites

Each method should be documented and provide a doc test (we are not giving examples here). In addition, any method defined for the objects or elements of a category should be supported by a test method, that is executed when running the test suite.

Base classes

In Sage, a "Parent" is an object of a category and contains elements. Parents should inherit from sage.structure.parent.Parent and their elements from sage.structure.element.

Sage provides appropriate sub-classes of Parent and Element for a variety of more concrete algebraic structures, such as groups, rings, or fields, and of their elements. But some old stuff in Sage doesn't use it. Volunteers for refactoring are welcome!

The parent Since we wish to implement a special kind of fields, namely fraction fields, it makes sense to build on top of the base class sage.rings.ring.Field provided by Sage.

```
sage: from sage.rings.ring import Field
```

This base class provides a lot more methods than a general parent:

```
sage: [p for p in dir(Field) if p not in dir(Parent)]
['__div__',
    '__fraction_field',
    '__ideal_monoid',
```

```
'___iter__',
'__pow__',
'__rdiv__',
'__rpow__',
'__rxor__',
'__xor__',
'_an_element',
'_an_element_c',
'_an_element_impl',
'_coerce_',
'_coerce_c',
'_coerce_impl',
'_coerce_try',
'_default_category',
'_gcd_univariate_polynomial',
_gens',
'_has_coerce_map_from',
'_ideal_class_',
'_latex_names',
'_list',
'_one_element',
'_pseudo_fraction_field',
'_random_nonzero_element',
'_unit_ideal',
^\prime_xgcd_univariate_polynomial^\prime,
'_zero_element',
'_zero_ideal',
'algebraic_closure',
'base_extend',
'cardinality',
'class_group',
'coerce_map_from_c',
'content',
'divides',
'epsilon',
'extension',
'fraction_field',
'frobenius_endomorphism',
'gcd',
'gen',
'gens',
'get_action_c',
'get_action_impl',
'has_coerce_map_from_c',
'ideal',
'ideal_monoid',
'integral_closure',
'is_commutative',
'is_field',
'is_finite',
'is_integral_domain',
'is_integrally_closed',
'is_noetherian',
'is_prime_field',
'is_ring',
'is_subring',
'krull_dimension',
'list',
```

```
'ngens',
'one',
'order',
'prime_subfield',
'principal_ideal',
'quo',
'quotient',
'quotient_ring',
'random_element',
'unit_ideal',
'zero',
'zero_ideal',
'zeta',
'zeta_order']
```

The following is a very basic implementation of fraction fields, that needs to be complemented later.

```
sage: from sage.structure.unique_representation import UniqueRepresentation
sage: class MyFrac(UniqueRepresentation, Field):
          def __init__(self, base):
              if base not in IntegralDomains():
. . . . :
                  raise ValueError, "%s is no integral domain"%base
. . . . :
              Field.__init__(self, base)
         def _repr_(self):
              return "NewFrac(%s)"%repr(self.base())
         def base_ring(self):
. . . . :
              return self.base().base_ring()
. . . . :
          def characteristic(self):
. . . . :
              return self.base().characteristic()
```

This basic implementation is formed by the following steps:

• Any ring in Sage has a **base** and a **base ring**. The "usual" fraction field of a ring R has the base R and the base ring R.base_ring():

```
sage: Frac(QQ['x']).base(), Frac(QQ['x']).base_ring()
(Univariate Polynomial Ring in x over Rational Field, Rational Field)
```

Declaring the base is easy: We just pass it as an argument to the field constructor.

```
\begin{tabular}{ll} \textbf{sage:} & Field(ZZ['x']).base() \\ & Univariate Polynomial Ring in x over Integer Ring \\ \end{tabular}
```

We are implementing a seperate method returning the base ring.

- Python uses double-underscore methods for arithmetic methods and string representations. Sage's base classes
 often have a default implementation, and it is requested to implement SINGLE underscore methods _repr_,
 and similarly _add_, _mul_ etc.
- You are encouraged to **make your parent "unique"**. That's to say, parents should only evaluate equal if they are identical. Sage provides frameworks to create unique parents. We use here the most easy one: Inheriting from the class sage.structure.unique_representation.UniqueRepresentation is enough. Making parents unique can be quite important for an efficient implementation, because the repeated creation of "the same" parent would take a lot of time.
- Fraction fields are only defined for integral domains. Hence, we raise an error if the given ring does not belong to the category of integral domains. This is our first use case of categories.
- Last, we add a method that returns the characteristic of the field. We don't go into details, but some automated tests that we study below implicitly rely on this method.

We see that our basic implementation correctly refuses a ring that is not an integral domain:

```
sage: MyFrac(ZZ['x'])
NewFrac(Univariate Polynomial Ring in x over Integer Ring)
sage: MyFrac(Integers(15))
Traceback (most recent call last):
...
ValueError: Ring of integers modulo 15 is no integral domain
```

Note: Inheritance from UniqueRepresentation automatically provides our class with pickling, preserving the unique parent condition. If we had defined the class in some external module or in an interactive session, pickling would work immediately.

However, for making the following example work in Sage's doctesting framework, we need to assign our class as an attribute of the __main__ module, so that the class can be looked up during unpickling.

```
sage: import __main__
sage: __main__.MyFrac = MyFrac
sage: loads(dumps(MyFrac(ZZ))) is MyFrac(ZZ)
True
```

Note: In the following sections, we will successively add or change details of MyFrac. Rather than giving a full class definition in each step, we define new versions of MyFrac by inheriting from the previously defined version of MyFrac. We believe this will help the reader to focus on the single detail that is relevant in each section.

The complete code can be found in the appendix.

The elements We use the base class sage.structure.element.FieldElement. Note that in the creation of field elements it is not tested that the given parent is a field:

```
sage: from sage.structure.element import FieldElement
sage: FieldElement(ZZ)
Generic element of a structure
```

Our toy implementation of fraction field elements is based on the following considerations:

- A fraction field element is defined by numerator and denominator, which both need to be elements of the base. There should be methods returning numerator resp. denominator.
- The denominator must not be zero, and (provided that the base is an ordered ring) we can make it non-negative, without loss of generality. By default, the denominator is one.
- The string representation is returned by the single-underscore method <u>repr</u>. In order to make our fraction field elements distinguishable from those already present in Sage, we use a different string representation.
- Arithmetic is implemented in single-underscore method _add_, _mul__, etc. We do not override the default double underscore __add__, _mul__, since otherwise, we could not use Sage's coercion model.
- Comparisons can be implemented using _cmp_. This automatically makes the relational operators like == and < work. In order to support the Python cmp() function, it is safest to define both _cmp_ and _cmp__ (because __cmp__ is not inherited if other comparison operators or __hash__ are defined). Of course you can just do __cmp__ = _cmp_.

Note that cmp should be provided, since otherwise comparison does not work:

```
sage: class Foo(sage.structure.element.Element):
....: def __init__(self, parent, x):
```

```
...: self.x = x
...: def _repr_(self):
...: return "<%s>"%self.x
sage: a = Foo(ZZ, 1)
sage: b = Foo(ZZ, 2)
sage: cmp(a,b)
Traceback (most recent call last):
...
NotImplementedError: comparison not implemented for <class '__main__.Foo'>
```

- In the single underscore methods, we can assume that *both arguments belong to the same parent*. This is one benefit of the coercion model.
- When constructing new elements as the result of arithmetic operations, we do not directly name our class, but we use self.__class__. Later, this will come in handy.

This gives rise to the following code:

```
sage: class MyElement (FieldElement):
           def __init__(self, parent, n, d=None):
                B = parent.base()
. . . . :
                if d is None:
. . . . :
                     d = B.one()
. . . . :
                if n not in B or d not in B:
. . . . :
                     raise ValueError("Numerator and denominator must be elements of %s"%B)
. . . . :
. . . . :
                # Numerator and denominator should not just be "in" B,
. . . . :
                # but should be defined as elements of B
                d = B(d)
. . . . :
                n = B(n)
. . . . :
                if d==0:
. . . . :
                     raise ZeroDivisionError("The denominator must not be zero")
. . . . :
                if d<0:
. . . . :
                     self.n = -n
                     self.d = -d
. . . . :
. . . . :
                else:
                     self.n = n
. . . . :
                     self.d = d
. . . . :
. . . . :
                FieldElement.__init__(self,parent)
. . . . :
           def numerator(self):
. . . . :
                return self.n
           def denominator(self):
. . . . :
                return self.d
. . . . :
           def _repr_(self):
. . . . :
                return "(%s):(%s)"%(self.n,self.d)
. . . . :
. . . . :
           def _cmp_(self, other):
                return cmp(self.n*other.denominator(), other.numerator()*self.d)
. . . . :
            \underline{\phantom{a}} cmp\underline{\phantom{a}} = \underline{\phantom{a}} cmp\underline{\phantom{a}}
           def _add_(self, other):
. . . . :
                C = self.__class_
. . . . :
                D = self.d*other.denominator()
. . . . :
                return C(self.parent(), self.n*other.denominator()+self.d*other.numerator(), D)
. . . . :
. . . . :
           def _sub_(self, other):
. . . . :
                C = self.__class_
                D = self.d*other.denominator()
. . . . :
                return C(self.parent(), self.n*other.denominator()-self.d*other.numerator(),D)
. . . . :
           def _mul_(self, other):
. . . . :
                C = self.__class__
. . . . :
                return C(self.parent(), self.n*other.numerator(), self.d*other.denominator())
. . . . :
           def _div_(self, other):
```

```
...: C = self.__class__
...: return C(self.parent(), self.n*other.denominator(), self.d*other.numerator())
```

Features and limitations of the basic implementation Thanks to the single underscore methods, some basic arithmetics works, **if** we stay inside a single parent structure:

```
sage: P = MyFrac(ZZ)
sage: a = MyElement(P, 3, 4)
sage: b = MyElement(P, 1, 2)
sage: a+b, a-b, a*b, a/b
((10):(8), (2):(8), (3):(8), (6):(4))
sage: a-b == MyElement(P, 1, 4)
True
```

We didn't implement exponentiation—but it just works:

```
sage: a^3
(27):(64)
```

There is a default implementation of element tests. We can already do

```
sage: a in P
True
```

since a is defined as an element of P. However, we can not verify yet that the integers are contained in the fraction field of the ring of integers. It does not even give a wrong answer, but results in an error:

```
sage: 1 in P
Traceback (most recent call last):
...
NotImplementedError
```

We will take care of this later.

Categories in Sage

Sometimes the base classes do not reflect the mathematics: The set of $m \times n$ matrices over a field forms, in general, not more than a vector space. Hence, this set (called MatrixSpace) is not implemented on top of sage.rings.ring.Ring. However, if m = n, then the matrix space is an algebra, thus, is a ring.

From the point of view of Python base classes, both cases are the same:

```
sage: MS1 = MatrixSpace(QQ,2,3)
sage: isinstance(MS1, Ring)
False
sage: MS2 = MatrixSpace(QQ,2)
sage: isinstance(MS2, Ring)
False
```

Sage's category framework can differentiate the two cases:

```
sage: Rings()
Category of rings
sage: MS1 in Rings()
False
```

```
sage: MS2 in Rings()
True
```

And indeed, MS2 has *more* methods than MS1:

```
sage: import inspect
sage: len([s for s in dir(MS1) if inspect.ismethod(getattr(MS1,s,None))])
59
sage: len([s for s in dir(MS2) if inspect.ismethod(getattr(MS2,s,None))])
87
```

This is because the class of MS2 also inherits from the parent class for algebras:

```
sage: MS1.__class__.__bases__
(<class 'sage.matrix.matrix_space.MatrixSpace'>,
<class 'sage.categories.category.JoinCategory.parent_class'>)
sage: MS2.__class__.__bases__
(<class 'sage.matrix.matrix_space.MatrixSpace'>,
<class 'sage.categories.category.JoinCategory.parent_class'>)
```

Below, we will explain how this can be taken advantage of.

It is no surprise that our parent P defined above knows that it belongs to the category of fields, as it is derived from the base class of fields.

```
sage: P.category()
Category of fields
```

However, we could choose a smaller category, namely the category of quotient fields.

Why should one choose a category? One can provide default methods for all objects of a category, and for all elements of such objects. Hence, the category framework is a way to inherit useful stuff that is not present in the base classes. These default methods do not rely on implementation details, but on mathematical concepts.

In addition, the categories define **test suites** for their objects and elements—see the last section. Hence, one also gets basic sanity tests for free.

How does the *category framework* **work?** Abstract base classes for the objects ("parent_class") and the elements of objects ("element_class") are provided by attributes of the category. During initialisation of a parent, the class of the parent is *dynamically changed* into a sub–class of the category's parent class. Likewise, sub–classes of the category's element class are available for the creation of elements of the parent, as explained below.

A dynamic change of classes does not work in Cython. Nevertheless, method inheritance still works, by virtue of a __getattr__ method.

Note: It is strongly recommended to use the category framework both in Python and in Cython.

Let us see whether there is any gain in chosing the category of quotient fields instead of the category of fields:

```
['_derivative', 'denominator', 'derivative', 'factor',
  'numerator', 'partial_fraction_decomposition']
```

So, there is no immediate gain for our fraction fields, but additional methods become available to our fraction field elements. Note that some of these methods are place-holders: There is no default implementation, but it is *required* (respectively is *optional*) to implement these methods:

```
sage: QuotientFields().element_class.denominator
<abstract method denominator at ...>
sage: from sage.misc.abstract_method import abstract_methods_of_class
sage: abstract_methods_of_class(QuotientFields().element_class)['optional']
['_add_', '_mul_']
sage: abstract_methods_of_class(QuotientFields().element_class)['required']
['__nonzero__', 'denominator', 'numerator']
```

Hence, when implementing elements of a quotient field, it is *required* to implement methods returning the denominator and the numerator, and a method that tells whether the element is nonzero, and in addition, it is *optional* (but certainly recommended) to provide some arithmetic methods. If one forgets to implement the required methods, the test suites of the category framework will complain—see below.

Implementing the category framework for the parent We simply need to declare the correct category by an optional argument of the field constructor, where we provide the possibility to override the default category:

When constructing instances of MyFrac, their class is dynamically changed into a new class called MyFrac_with_category. It is a common sub-class of MyFrac and of the category's parent class:

```
sage: P = MyFrac(ZZ)
sage: type(P)
<class '__main__.MyFrac_with_category'>
sage: isinstance(P,MyFrac)
True
sage: isinstance(P,QuotientFields().parent_class)
True
```

The fraction field P inherits additional methods. For example, the base class Field does not have a method sum. But P inherits such method from the category of commutative additive monoids—see sum ():

```
sage: P.sum.__module__
'sage.categories.additive_monoids'
```

We have seen above that we can add elements. Nevertheless, the sum method does not work, yet:

```
sage: a = MyElement(P, 3, 4)
sage: b = MyElement(P, 1, 2)
sage: c = MyElement(P, -1, 2)
sage: P.sum([a, b, c])
Traceback (most recent call last):
...
NotImplementedError
```

The reason is that the sum method starts with the return value of P.zero(), which defaults to P(0)—but the conversion of integers into P is not implemented, yet.

Implementing the category framework for the elements Similar to what we have seen for parents, a new class is dynamically created that combines the element class of the parent's category with the class that we have implemented above. However, the category framework is implemented in a different way for elements than for parents:

- We provide the parent P (or its class) with an attribute called "Element", whose value is a class.
- The parent *automatically* obtains an attribute P.element_class, that subclasses both P.Element and P.category().element_class.

Hence, for providing our fraction fields with their own element classes, we just need to add a single line to our class:

```
sage: class MyFrac(MyFrac):
....: Element = MyElement
```

This little change provides several benefits:

• We can now create elements by simply calling the parent:

```
sage: P = MyFrac(ZZ)
sage: P(1), P(2,3)
((1):(1), (2):(3))
```

• There is a method zero returning the expected result:

```
sage: P.zero()
(0):(1)
```

• The sum method mentioned above suddenly works:

```
sage: a = MyElement(P, 9, 4)
sage: b = MyElement(P, 1, 2)
sage: c = MyElement(P, -1, 2)
sage: P.sum([a,b,c])
(36):(16)
```

What did happen behind the scenes to make this work? We provided P.Element, and thus obtain P.element_class, which is a *lazy attribute*. It provides a *dynamic* class, which is a sub-class of both MyElement defined above and of P.category().element_class:

```
sage: P.__class__.element_class
<sage.misc.lazy_attribute.lazy_attribute object at ...>
sage: P.element_class
<class '__main__.MyFrac_with_category.element_class'>
sage: type(P.element_class)
<class 'sage.structure.dynamic_class.DynamicInheritComparisonMetaclass'>
sage: issubclass(P.element_class, MyElement)
True
sage: issubclass(P.element_class, P.category().element_class)
True
```

The default __call__ method of P passes the given arguments to P.element_class, adding the argument parent=P. This is why we are now able to create elements by calling the parent.

In particular, these elements are instances of that new dynamic class:

```
sage: type(P(2,3))
<class '__main__.MyFrac_with_category.element_class'>
```

Note: All elements of P should use the element class. In order to make sure that this also holds for the result of arithmetic operations, we created them as instances of self. __class__ in the arithmetic methods of MyElement.

P.zero() defaults to returning P(0) and thus returns an instance of P.element_class. Since P.sum([...]) starts the summation with P.zero() and the class of the sum only depends on the first summand, by our implementation, we have:

```
sage: type(a)
<class '__main__.MyElement'>
sage: isinstance(a,P.element_class)
False
sage: type(P.sum([a,b,c]))
<class '__main__.MyFrac_with_category.element_class'>
```

The method factor provided by P.category().element_class(see above) simply works:

```
sage: a; a.factor(); P(6,4).factor()
(9):(4)
2^-2 * 3^2
2^-1 * 3
```

But that's surprising: The element a is just an instance of MyElement, but not of P.element_class, and its class does not know about the factor method. In fact, this is due to a __getattr__ method defined for sage.structure.element.Element.

```
sage: hasattr(type(a), 'factor')
False
sage: hasattr(P.element_class, 'factor')
True
sage: hasattr(a, 'factor')
True
```

A first note on performance The category framework is sometimes blamed for speed regressions, as in trac ticket #9138 and trac ticket #11900. But if the category framework is *used properly*, then it is fast. For illustration, we determine the time needed to access an attribute inherited from the element class. First, we consider an element that uses the class that we implemented above, but does not use the category framework properly:

```
sage: type(a)
<class '__main__.MyElement'>
sage: timeit('a.factor',number=1000) # random
1000 loops, best of 3: 2 us per loop
```

Now, we consider an element that is equal to a, but uses the category framework properly:

```
sage: a2 = P(9,4)
sage: a2 == a
True
sage: type(a2)
<class '__main__.MyFrac_with_category.element_class'>
sage: timeit('a2.factor',number=1000) # random
1000 loops, best of 3: 365 ns per loop
```

So, don't be afraid of using categories!

Coercion—the basics

Theoretical background

Coercion is not just *type conversion* "Coercion" in the C programming language means "automatic type conversion". However, in Sage, coercion is involved if one wants to be able to do arithmetic, comparisons, etc. between elements of distinct parents. Hence, **coercion is not about a change of types, but about a change of parents.**

As an illustration, we show that elements of the same type may very well belong to rather different parents:

```
sage: P1 = QQ['v,w']; P2 = ZZ['w,v']
sage: type(P1.gen()) == type(P2.gen())
True
sage: P1 == P2
False
```

 P_2 naturally is a sub-ring of P_1 . So, it makes sense to be able to add elements of the two rings—the result should then live in P_1 , and indeed it does:

```
sage: (P1.gen()+P2.gen()).parent() is P1
True
```

It would be rather inconvenient if one needed to *manually* convert an element of P_2 into P_1 before adding. The coercion system does that conversion automatically.

Not every conversion is a coercion A coercion happens implicitly, without being explicitly requested by the user. Hence, coercion must be based on mathematical rigour. In our example, any element of P_2 can be naturally interpreted as an element of P_1 . We thus have:

```
sage: P1.has_coerce_map_from(P2)
True
sage: P1.coerce_map_from(P2)
Conversion map:
   From: Multivariate Polynomial Ring in w, v over Integer Ring
   To: Multivariate Polynomial Ring in v, w over Rational Field
```

While there is a conversion from P_1 to P_2 (namely restricted to polynomials with integral coefficients), this conversion is not a coercion:

```
sage: P2.convert_map_from(P1)
Conversion map:
   From: Multivariate Polynomial Ring in v, w over Rational Field
   To: Multivariate Polynomial Ring in w, v over Integer Ring
sage: P2.has_coerce_map_from(P1)
False
sage: P2.coerce_map_from(P1) is None
True
```

The four axioms requested for coercions

1. A coercion is a morphism in an appropriate category.

This first axiom has two implications:

(a) A coercion is defined on all elements of a parent.

A polynomial of degree zero over the integers can be interpreted as an integer—but the attempt to convert a polynomial of non-zero degree would result in an error:

```
sage: ZZ(P2.one())
1
sage: ZZ(P2.gen(1))
Traceback (most recent call last):
...
TypeError: not a constant polynomial
```

Hence, we only have a *partial* map. This is fine for a *conversion*, but a partial map does not qualify as a *coercion*.

(b) Coercions are structure preserving.

Any real number can be converted to an integer, namely by rounding. However, such a conversion is not useful in arithmetic operations, since the underlying algebraic structure is not preserved:

```
sage: int(1.6)+int(2.7) == int(1.6+2.7)
False
```

The structure that is to be preserved depends on the category of the involved parents. For example, the coercion from the integers into the rational field is a homomorphism of euclidean domains:

```
sage: QQ.coerce_map_from(ZZ).category_for()
Join of Category of euclidean domains and Category of metric spaces
```

2. There is at most one coercion from one parent to another

In addition, if there is a *coercion* from P_2 to P_1 , then a *conversion* from P_2 to P_1 is defined for all elements of P_2 and coincides with the coercion. Nonetheless, user-exposed maps are copies of the internally used maps whence the lack of identity between different instantiations:

```
sage: P1.coerce_map_from(P2) is P1.convert_map_from(P2)
False
```

For internally used maps, the maps are identical:

```
sage: P1._internal_coerce_map_from(P2) is P1._internal_convert_map_from(P2)
True
```

3. Coercions can be composed

If there is a coercion $\varphi: P_1 \to P_2$ and another coercion $\psi: P_2 \to P_3$, then the composition of φ followed by ψ must yield the unique coercion from P_1 to P_3 .

4. The identity is a coercion

Together with the two preceding axioms, it follows: If there are coercions from P_1 to P_2 and from P_2 to P_1 , then they are mutually inverse.

Implementing a conversion We have seen above that some conversions into our fraction fields became available after providing the attribute Element. However, we can not convert elements of a fraction field into elements of another fraction field, yet:

```
sage: P(2/3)
Traceback (most recent call last):
...
ValueError: Numerator and denominator must be elements of Integer Ring
```

For implementing a conversion, the default __call__ method should (almost) never be overridden. Instead, we implement the method _element_constructor_, that should return an instance of the parent's element class. Some old parent classes violate that rule—please help to refactor them!

```
sage: class MyFrac(MyFrac):
           def _element_constructor_(self, *args, **kwds):
. . . . :
               if len(args)!=1:
. . . . :
                    return self.element_class(self, *args, **kwds)
. . . . :
. . . . :
               x = args[0]
. . . . :
               try:
                    P = x.parent()
. . . . :
               except AttributeError:
. . . . :
                   return self.element_class(self, x, **kwds)
. . . . :
               if P in QuotientFields() and P != self.base():
. . . . :
                   return self.element_class(self, x.numerator(), x.denominator(), **kwds)
. . . . :
               return self.element_class(self, x, **kwds)
. . . . :
```

In addition to the conversion from the base ring and from pairs of base ring elements, we now also have a conversion from the rationals to our fraction field of **Z**:

```
sage: P = MyFrac(ZZ)
sage: P(2); P(2,3); P(3/4)
(2):(1)
(2):(3)
(3):(4)
```

Recall that above, the test $1 \in P$ failed with an error. We try again and find that the error has disappeared. This is because we are now able to convert the integer 1 into P. But the containment test still yields a wrong answer:

```
sage: 1 in P
False
```

The technical reason: We have a conversion P(1) of 1 into P, but this is not known as a coercion—yet!

```
sage: P.has_coerce_map_from(ZZ), P.has_coerce_map_from(QQ)
(False, False)
```

Establishing a coercion There are two main ways to make Sage use a particular conversion as a coercion:

- One can use sage.structure.parent.Parent.register_coercion(), normally during initialisation of the parent (see documentation of the method).
- A more flexible way is to provide a method coerce map from for the parent.

Let P and R be parents. If P._coerce_map_from_(R) returns False or None, then there is no coercion from R to P. If it returns a map with domain R and codomain P, then this map is used for coercion. If it returns True, then the conversion from R to P is used as coercion.

Note that in the following implementation, we need a special case for the rational field, since QQ.base() is not the ring of integers.

```
....: if hasattr(S,'ring_of_integers') and self.base().has_coerce_map_from(S.ring_of_integers')
....:
```

By the method above, a parent coercing into the base ring will also coerce into the fraction field, and a fraction field coerces into another fraction field if there is a coercion of the corresponding base rings. Now, we have:

```
sage: P = MyFrac(QQ['x'])
sage: P.has_coerce_map_from(ZZ['x']), P.has_coerce_map_from(Frac(ZZ['x'])), P.has_coerce_map_from(QQ
(True, True, True)
```

We can now use coercion from $\mathbb{Z}[x]$ and from \mathbb{Q} into P for arithmetic operations between the two rings:

```
sage: 3/4+P(2)+ZZ['x'].gen(), (P(2)+ZZ['x'].gen()).parent() is P(4*x + 11):(4), True)
```

Equality and element containment Recall that above, the test $1 \in P$ gave a wrong answer. Let us repeat the test now:

```
sage: 1 in P
True
```

Why is that?

The default element containment test $x \in P$ is based on the interplay of three building blocks: conversion, coercion, and equality test.

- 1. Clearly, if the conversion P(x) raises an error, then x can not be seen as an element of P. On the other hand, a conversion P(x) can generally do very nasty things. So, the fact that P(x) works without error is necessary, but not sufficient for $x \in P$.
- 2. If P is the parent of x, then the conversion P(x) will not change x (at least, that's the default). Hence, we will have x = P(x).
- 3. Sage uses coercion not only for arithmetic operations, but also for comparison: *If* there is a coercion from the parent of x to P, then the equality test x==P(x) reduces to P(x)==P(x). Otherwise, x==P(x) will evaluate as false.

That leads to the following default implementation of element containment testing:

```
Note: x \in P holds if and only if the test x = P(x) does not raise an error and evaluates as true.
```

If the user is not happy with that behaviour, the "magical" Python method __contains__ can be overridden.

Coercion—the advanced parts

So far, we are able to add integers and rational numbers to elements of our new implementation of the fraction field of **Z**.

```
sage: P = MyFrac(ZZ)
sage: 1/2+P(2,3)+1
(13):(6)
```

Surprisingly, we can even add a polynomial over the integers to an element of P, even though the *result lives in a new parent*, namely in a polynomial ring over P:

```
sage: P(1/2) + ZZ['x'].gen(), (P(1/2) + ZZ['x'].gen()).parent() is P['x'] ((1):(1)*x + (1):(2), True)
```

In the next, seemingly more easy example, there "obviously" is a coercion from the fraction field of \mathbf{Z} to the fraction field of $\mathbf{Z}[x]$. However, Sage does not know enough about our new implementation of fraction fields. Hence, it does not recognise the coercion:

```
sage: Frac(ZZ['x']).has_coerce_map_from(P)
False
```

Two obvious questions arise:

- 1. How / why has the new ring been constructed in the example above?
- 2. How can we establish a coercion from P to $Frac(\mathbf{Z}[x])$?

The key to answering both question is the construction of parents from simpler pieces, that we are studying now. Note that we will answer the second question *not* by providing a coercion from P to $\operatorname{Frac}(\mathbf{Z}[x])$, but by teaching Sage to automatically construct $\operatorname{MyFrac}(\mathbf{Z}[x])$ and coerce both P and $\operatorname{Frac}(\mathbf{Z}[x])$ into it.

If we are lucky, a parent can tell how it has been constructed:

```
sage: Poly,R = QQ['x'].construction()
sage: Poly,R
(Poly[x], Rational Field)
sage: Fract,R = QQ.construction()
sage: Fract,R
(FractionField, Integer Ring)
```

In both cases, the first value returned by construction() is a mathematical construction, called *construction functor*—see ConstructionFunctor. The second return value is a simpler parent to which the construction functor is applied.

Being functors, the same construction can be applied to different objects of a category:

```
sage: Poly(QQ) is QQ['x']
True
sage: Poly(ZZ) is ZZ['x']
True
sage: Poly(P) is P['x']
True
sage: Fract(QQ['x'])
Fraction Field of Univariate Polynomial Ring in x over Rational Field
```

Let us see on which categories these construction functors are defined:

```
sage: Poly.domain()
Category of rings
sage: Poly.codomain()
Category of rings
sage: Fract.domain()
Category of integral domains
sage: Fract.codomain()
Category of fields
```

In particular, the construction functors can be composed:

```
sage: Poly*Fract
Poly[x](FractionField(...))
```

```
sage: (Poly*Fract)(ZZ) is QQ['x']
True
```

In addition, it is often assumed that we have a coercion from input to output of the construction functor:

```
sage: ((Poly*Fract)(ZZ)).coerce_map_from(ZZ)
Composite map:
    From: Integer Ring
    To: Univariate Polynomial Ring in x over Rational Field
    Defn: Natural morphism:
        From: Integer Ring
        To: Rational Field
        then
            Polynomial base injection morphism:
        From: Rational Field
        To: Univariate Polynomial Ring in x over Rational Field
```

Construction functors do not necessarily commute:

```
sage: (Fract*Poly)(ZZ)
Fraction Field of Univariate Polynomial Ring in x over Integer Ring
```

The pushout of construction functors We can now formulate our problem. We have parents P_1 , P_2 and R, and construction functors F_1 , F_2 , such that $P_1 = F_1(R)$ and $P_2 = F_2(R)$. We want to find a new construction functor F_3 , such that both P_1 and P_2 coerce into $P_3 = F_3(R)$.

In analogy to a notion of category theory, P_3 is called the pushout () of P_1 and P_2 ; and similarly F_3 is called the pushout of F_1 and F_2 .

```
sage: from sage.categories.pushout import pushout
sage: pushout(Fract(ZZ),Poly(ZZ))
Univariate Polynomial Ring in x over Rational Field
```

 $F_1 \circ F_2$ and $F_2 \circ F_1$ are natural candidates for the pushout of F_1 and F_2 . However, the order of the functors must rely on a canonical choice. "Indecomposable" construction functors have a *rank*, and this allows to order them canonically:

Note: If F1. rank is smaller than F2. rank, then the pushout is $F_2 \circ F_1$ (hence, F_1 is applied first).

We have

```
sage: Fract.rank, Poly.rank
(5, 9)

and thus the pushout is

sage: Fract.pushout(Poly), Poly.pushout(Fract)
(Poly[x](FractionField(...)), Poly[x](FractionField(...)))
```

This is why the example above has worked.

However, only "elementary" construction functors have a rank:

```
sage: (Fract*Poly).rank
Traceback (most recent call last):
...
AttributeError: 'CompositeConstructionFunctor' object has no attribute 'rank'
```

Shuffling composite construction functors If composed construction fuctors ... $\circ F_2 \circ F_1$ and ... $\circ G_2 \circ G_1$ are given, then Sage determines their pushout by *shuffling* the constituents:

- If F1.rank < G1.rank then we apply F_1 first, and continue with ... $\circ F_3 \circ F_2$ and ... $\circ G_2 \circ G_1$.
- If F1.rank > G1.rank then we apply G_1 first, and continue with ... $\circ F_2 \circ F_1$ and ... $\circ G_3 \circ G_2$.

If F1.rank == G1.rank, then the tie needs to be broken by other techniques (see below).

As an illustration, we first get us some functors and then see how chains of functors are shuffled.

```
sage: AlgClos, R = CC.construction(); AlgClos
AlgebraicClosureFunctor

sage: Compl, R = RR.construction(); Compl
Completion[+Infinity]

sage: Matr, R = (MatrixSpace(ZZ,3)).construction(); Matr
MatrixFunctor

sage: AlgClos.rank, Compl.rank, Fract.rank, Poly.rank, Matr.rank
(3, 4, 5, 9, 10)
```

When we apply Fract, AlgClos, Poly and Fract to the ring of integers, we obtain:

```
sage: (Fract*Poly*AlgClos*Fract)(ZZ)
Fraction Field of Univariate Polynomial Ring in x over Algebraic Field
```

When we apply Compl, Matr and Poly to the ring of integers, we obtain:

```
sage: (Poly*Matr*Compl)(ZZ)
Univariate Polynomial Ring in x over Full MatrixSpace of 3 by 3 dense matrices over Real Field with
```

Applying the shuffling procedure yields

```
sage: (Poly*Matr*Fract*Poly*AlgClos*Fract*Compl)(ZZ)
Univariate Polynomial Ring in x over Full MatrixSpace of 3 by 3 dense matrices over Fraction Field or
```

and this is indeed equal to the pushout found by Sage:

```
sage: pushout((Fract*Poly*AlgClos*Fract)(ZZ), (Poly*Matr*Compl)(ZZ))
Univariate Polynomial Ring in x over Full MatrixSpace of 3 by 3 dense matrices over Fraction Field or
```

Breaking the tie If F1.rank==G1.rank then Sage's pushout constructions offers two ways to proceed:

- 1. Construction functors have a method merge () that either returns None or returns a construction functor—see below. If either F1.merge (G1) or G1.merge (F1) returns a construction functor H_1 , then we apply H_1 and continue with ... $\circ F_3 \circ F_2$ and ... $\circ G_3 \circ G_2$.
- 2. Construction functors have a method commutes (). If either F1.commutes (G1) or G1.commutes (F1) returns True, then we apply both F_1 and G_1 in any order, and continue with ... $\circ F_3 \circ F_2$ and ... $\circ G_3 \circ G_2$.

By default, F1.merge (G1) returns F1 if F1==G1, and returns None otherwise. The commutes () method exists, but it seems that so far nobody has implemented two functors of the same rank that commute.

Establishing a default implementation The typical application of merge() is to provide a coercion between *different implementations* of the *same algebraic structure*.

Note: If F1(P) and F2(P) are different implementations of the same thing, then F1.merge(F2)(P) should return the default implementation.

We want to boldly turn our toy implementation of fraction fields into the new default implementation. Hence:

- Next, we implement a new version of the "usual" fraction field functor, having the same rank, but returning our new implementation.
- We make our new implementation the default, by virtue of a merge method.

Warning:

- Do not override the default __call__ method of ConstructionFunctor—implement _apply_functor instead.
- Declare domain and codomain of the functor during initialisation.

```
sage: from sage.categories.pushout import ConstructionFunctor
sage: class MyFracFunctor(ConstructionFunctor):
         rank = 5
          def __init__(self):
. . . . :
              ConstructionFunctor.__init__(self, IntegralDomains(), Fields())
          def _apply_functor(self, R):
              return MyFrac(R)
. . . . :
         def merge(self, other):
              if isinstance(other, (type(self), sage.categories.pushout.FractionField)):
. . . . :
                  return self
. . . . :
sage: MyFracFunctor()
MyFracFunctor
```

We verify that our functor can really be used to construct our implementation of fraction fields, and that it can be merged with either itself or the usual fraction field constructor:

```
sage: MyFracFunctor() (ZZ)
NewFrac(Integer Ring)

sage: MyFracFunctor().merge(MyFracFunctor())
MyFracFunctor

sage: MyFracFunctor().merge(Fract)
MyFracFunctor
```

There remains to let our new fraction fields know about the new construction functor:

```
sage: class MyFrac(MyFrac):
....:     def construction(self):
....:     return MyFracFunctor(), self.base()

sage: MyFrac(ZZ['x']).construction()
(MyFracFunctor, Univariate Polynomial Ring in x over Integer Ring)
```

Due to merging, we have:

```
sage: pushout(MyFrac(ZZ['x']), Frac(QQ['x']))
NewFrac(Univariate Polynomial Ring in x over Rational Field)
```

A second note on performance Being able to do arithmetics involving elements of different parents, with the automatic creation of a pushout to contain the result, is certainly convenient—but one should not rely on it, if speed matters. Simply the conversion of elements into different parents takes time. Moreover, by trac ticket #14058, the pushout may be subject to Python's cyclic garbage collection. Hence, if one does not keep a strong reference to it, the same parent may be created repeatedly, which is a waste of time. In the following example, we illustrate the slow–down resulting from blindly relying on coercion:

```
sage: ZZxy = ZZ['x','y']
sage: a = ZZxy('x')
sage: b = 1/2
sage: timeit("c = a+b")  # random
10000 loops, best of 3: 172 us per loop
sage: QQxy = QQ['x','y']
sage: timeit("c2 = QQxy(a)+QQxy(b)") # random
10000 loops, best of 3: 168 us per loop
sage: a2 = QQxy(a)
sage: b2 = QQxy(b)
sage: timeit("c2 = a2+b2") # random
100000 loops, best of 3: 10.5 us per loop
```

Hence, if one avoids the explicit or implicit conversion into the pushout, but works in the pushout right away, one can get a more than 10–fold speed–up.

The test suites of the category framework

The category framework does not only provide functionality but also a test framework. This section logically belongs to the section on categories, but without the bits that we have implemented in the section on coercion, our implementation of fraction fields would not have passed the tests yet.

"Abstract" methods We have already seen above that a category can require/suggest certain parent or element methods, that the user must/should implement. This is in order to smoothly blend with the methods that already exist in Sage.

The methods that ought to be provided are called abstract_method(). Let us see what methods are needed for quotient fields and their elements:

```
sage: from sage.misc.abstract_method import abstract_methods_of_class
sage: abstract_methods_of_class(QuotientFields().parent_class)['optional']
[]
sage: abstract_methods_of_class(QuotientFields().parent_class)['required']
['__contains__']
```

Hence, the only required method (that is actually required for all parents that belong to the category of sets) is an element containment test. That's fine, because the base class Parent provides a default containment test.

The elements have to provide more:

```
sage: abstract_methods_of_class(QuotientFields().element_class)['optional']
['_add_', '_mul_']
```

```
sage: abstract_methods_of_class(QuotientFields().element_class)['required']
['__nonzero__', 'denominator', 'numerator']
```

Hence, the elements must provide denominator() and numerator() methods, and must be able to tell whether they are zero or not. The base class Element provides a default __nonzero__() method. In addition, the elements may provide Sage's single underscore arithmetic methods (actually any ring element *should* provide them).

The _test_... methods If a parent or element method's name start with "_test_", it gives rise to a test in the automatic test suite. For example, it is tested

- whether a parent P actually is an instance of the parent class of the category of P,
- whether the user has implemented the required abstract methods,
- whether some defining structural properties (e.g., commutativity) hold.

For example, if one forgets to implement required methods, one obtains the following error:

```
sage: class Foo(Parent):
....: Element = sage.structure.element.Element
....: def __init__(self):
....: Parent.__init__(self, category=QuotientFields())
sage: Bar = Foo()
sage: bar = Bar.element_class(Bar)
sage: bar._test_not_implemented_methods()
Traceback (most recent call last):
...
AssertionError: Not implemented method: denominator
```

Here are the tests that form the test suite of quotient fields:

```
sage: [t for t in dir(QuotientFields().parent_class) if t.startswith('_test_')]
['_test_additive_associativity',
  _test_an_element',
 '_test_associativity',
 '_test_cardinality',
 '_test_characteristic',
 '_test_characteristic_fields',
 '_test_distributivity',
 '_test_elements',
 '_test_elements_eq_reflexive',
 '_test_elements_eq_symmetric',
 '_test_elements_eq_transitive',
 '_test_elements_neq',
 '_test_euclidean_degree',
 '_test_gcd_vs_xgcd',
 '_test_one', '_test_prod',
 '_test_quo_rem',
 '_test_some_elements',
 '_test_zero',
 '_test_zero_divisors']
```

We have implemented all abstract methods (or inherit them from base classes), we use the category framework, and we have implemented coercions. So, we are confident that the test suite runs without an error. In fact, it does!

Note: The following trick with the __main__ module is only needed in doctests, not in an interactive session or when defining the classes externally.

```
sage: __main__.MyFrac = MyFrac
sage: __main__.MyElement = MyElement
sage: P = MyFrac(ZZ['x'])
sage: TestSuite(P).run()
```

Let us see what tests are actually performed:

```
sage: TestSuite(P).run(verbose=True)
running ._test_additive_associativity() . . . pass
running ._test_an_element() . . . pass
running ._test_associativity() . . . pass
running ._test_cardinality() . . . pass
running ._test_category() . . . pass
running ._test_characteristic() . . . pass
running ._test_characteristic_fields() . . . pass
running ._test_distributivity() . . . pass
running ._test_elements() . . .
 Running the test suite of self.an_element()
 running ._test_category() . . . pass
 running ._test_eq() . . . pass
 running ._test_nonzero_equal() . . . pass
 running ._test_not_implemented_methods() . . . pass
 running ._test_pickling() . . . pass
 pass
running ._test_elements_eq_reflexive() . . . pass
running ._test_elements_eq_symmetric() . . . pass
running ._test_elements_eq_transitive() . . . pass
running ._test_elements_neq() . . . pass
running ._test_eq() . . . pass
running ._test_euclidean_degree() . . . pass
running ._test_gcd_vs_xgcd() . . . pass
running ._test_not_implemented_methods() . . . pass
running ._test_one() . . . pass
running ._test_pickling() . . . pass
running ._test_prod() . . . pass
running ._test_quo_rem() . . . pass
running ._test_some_elements() . . . pass
running ._test_zero() . . . pass
running ._test_zero_divisors() . . . pass
```

Implementing a new category with additional tests As one can see, tests are also performed on elements. There are methods that return one element or a list of some elements, relying on "typical" elements that can be found in most algebraic structures.

```
sage: P.an_element(); P.some_elements()
(2):(1)
[(2):(1)]
```

Unfortunately, the list of elements that is returned by the default method is of length one, and that single element could also be a bit more interesting. The method an_element relies on a method _an_element_(), so, we implement that. We also override the some_elements method.

```
sage: class MyFrac(MyFrac):
....:     def _an_element_(self):
....:     a = self.base().an_element()
....:     b = self.base_ring().an_element()
```

```
if (a+b)!=0:
                   return self(a)**2/(self(a+b)**3)
. . . . :
               if b != 0:
. . . . :
                   return self(a)/self(b)**2
. . . . :
               return self(a) **2*self(b) **3
. . . . :
          def some_elements(self):
. . . . :
               return [self.an_element(), self(self.base().an_element()), self(self.base_ring().an_element())
sage: P = MyFrac(ZZ['x'])
sage: P.an_element(); P.some_elements()
(x^2): (x^3 + 3*x^2 + 3*x + 1)
[(x^2):(x^3 + 3*x^2 + 3*x + 1), (x):(1), (1):(1)]
```

Now, as we have more interesting elements, we may also add a test for the "factor" method. Recall that the method was inherited from the category, but it appears that it is not tested.

Normally, a test for a method defined by a category should be provided by the same category. Hence, since factor is defined in the category of quotient fields, a test should be added there. But we won't change source code here and will instead create a sub-category.

Apparently, If e is an element of a quotient field, the product of the factors returned by e. factor () should be equal to e. For forming the product, we use the prod method, that, no surprise, is inherited from another category:

```
sage: P.prod.__module__
'sage.categories.monoids'
```

When we want to create a sub-category, we need to provide a method super_categories (), that returns a list of all immediate super categories (here: category of quotient fields).

```
Warning: A sub-category S of a category C is not implemented as a sub-class of C.__class__! S becomes a sub-category of C only if S.super_categories () returns (a sub-category of) C!
```

The parent and element methods of a category are provided as methods of classes that are the attributes ParentMethods and Element Methods of the category, as follows:

```
sage: from sage.categories.category import Category
sage: class QuotientFieldsWithTest(Category): # do *not* inherit from QuotientFields, but ...
         def super_categories(self):
. . . . :
              return [QuotientFields()]
                                                # ... declare QuotientFields as a super category!
          class ParentMethods:
. . . . :
              pass
. . . . :
          class ElementMethods:
. . . . :
              def _test_factorisation(self, **options):
                  P = self.parent()
                  assert self == P.prod([P(b)**e for b,e in self.factor()])
. . . . :
```

We provide an instance of our quotient field implementation with that new category. Note that categories have a default _repr_ method, that guesses a good string representation from the name of the class: QuotientFieldsWithTest becomes "quotient fields with test".

Note: The following trick with the __main__ module is only needed in doctests, not in an interactive session or when defining the classes externally.

```
sage: __main__.MyFrac = MyFrac
sage: __main__.MyElement = MyElement
sage: __main__.QuotientFieldsWithTest = QuotientFieldsWithTest
```

Last, we observe that the new test has automatically become part of the test suite. We remark that the existing tests became more serious as well, since we made sage.structure.parent.Parent.an_element() return something more interesting.

```
sage: TestSuite(P).run(verbose=True)
running ._test_additive_associativity() . . . pass
running ._test_an_element() . . . pass
running ._test_associativity() . . . pass
running ._test_cardinality() . . . pass
running ._test_category() . . . pass
running ._test_characteristic() . . . pass
running ._test_characteristic_fields() . . . pass
running ._test_distributivity() . . . pass
running ._test_elements() . . .
 Running the test suite of self.an_element()
 running ._test_category() . . . pass
 running ._test_eq() . . . pass
 running ._test_factorisation() . . . pass
 running ._test_nonzero_equal() . . . pass
 running ._test_not_implemented_methods() . . . pass
 running ._test_pickling() . . . pass
 pass
running ._test_elements_eq_reflexive() . . . pass
running ._test_elements_eq_symmetric() . . . pass
running ._test_elements_eq_transitive() . . . pass
running ._test_elements_neq() . . . pass
running ._test_eq() . . . pass
running ._test_euclidean_degree() . . . pass
running ._test_gcd_vs_xgcd() . . . pass
running ._test_not_implemented_methods() . . . pass
running ._test_one() . . . pass
running ._test_pickling() . . . pass
running ._test_prod() . . . pass
running ._test_quo_rem() . . . pass
running ._test_some_elements() . . . pass
running ._test_zero() . . . pass
running ._test_zero_divisors() . . . pass
```

Appendix: The complete code

```
# Importing base classes, ...
import sage
```

```
from sage.rings.ring import Field
   from sage.structure.element import FieldElement
   from sage.categories.category import Category
   # ... the UniqueRepresentation tool,
   from sage.structure.unique_representation import UniqueRepresentation
   # ... some categories, and ...
   from sage.categories.fields import Fields
   from sage.categories.quotient_fields import QuotientFields
   from sage.categories.integral_domains import IntegralDomains
   # construction functors
   from sage.categories.pushout import ConstructionFunctor
14
   # Fraction field elements
15
   class MyElement (FieldElement):
16
       def __init__(self, parent, n, d=None):
17
           if parent is None:
18
                raise ValueError("The parent must be provided")
19
           B = parent.base()
20
           if d is None:
21
                # The default denominator is one
22
                d = B.one()
23
            # verify that both numerator and denominator belong to the base
24
           if n not in B or d not in B:
                raise ValueError("Numerator and denominator must be elements of %s"%B)
26
            # Numerator and denominator should not just be "in" B,
27
            # but should be defined as elements of B
28
           d = B(d)
29
           n = B(n)
30
            # the denominator must not be zero
31
           if d==0:
32
33
                raise ZeroDivisionError("The denominator must not be zero")
            # normalize the denominator: WLOG, it shall be non-negative.
34
           if d<0:
35
                self.n = -n
36
                self.d = -d
37
           else:
38
                self.n = n
                self.d = d
40
           FieldElement.__init__(self,parent)
41
42.
       # Methods required by the category of fraction fields:
43
       def numerator(self):
44
           return self.n
46
       def denominator(self):
47
           return self.d
48
       # String representation (single underscore!)
49
       def _repr_(self):
50
           return "(%s):(%s)"%(self.n, self.d)
51
       # Comparison: We can assume that both arguments are coerced
53
        # into the same parent, which is a fraction field. Hence, we
54
       # are allowed to use the denominator() and numerator() methods
55
       # on the second argument.
56
       def _cmp_(self, other):
57
            return cmp(self.n*other.denominator(), other.numerator()*self.d)
        # Support for cmp() (in this example, we don't define __hash__
```

```
# so this is not strictly needed)
62
        \underline{\hspace{0.1cm}} cmp\underline{\hspace{0.1cm}} = \underline{\hspace{0.1cm}} cmp\underline{\hspace{0.1cm}}
63
        # Arithmetic methods, single underscore. We can assume that both
        # arguments are coerced into the same parent.
65
        # We return instances of self.__class__, because self.__class__ will
66
        # eventually be a sub-class of MyElement.
67
        def _add_(self, other):
68
            C = self.__class__
69
            D = self.d*other.denominator()
70
            return C(self.parent(), self.n*other.denominator()+self.d*other.numerator(),D)
71
        def _sub_(self, other):
72
            C = self.__class__
73
            D = self.d*other.denominator()
74
            return C(self.parent(), self.n*other.denominator()-self.d*other.numerator(),D)
75
        def _mul_(self, other):
76
            C = self.__class___
77
            return C(self.parent(), self.n*other.numerator(), self.d*other.denominator())
78
        def _div_(self, other):
79
            C = self.__class__
80
            return C(self.parent(), self.n*other.denominator(), self.d*other.numerator())
81
82
    # Inheritance from UniqueRepresentation implements the unique parent
83
    # behaviour. Moreover, it implements pickling (provided that Python
    # succeeds to look up the class definition).
    class MyFrac (UniqueRepresentation, Field):
86
        # Implement the category framework for elements, which also
87
        # makes some basic conversions work.
88
        Element = MyElement
89
91
        # Allow to pass to a different category, by an optional argument
        def __init__(self, base, category=None):
92
             # Fraction fields only exist for integral domains
93
            if base not in IntegralDomains():
94
                 raise ValueError, "%s is no integral domain"%base
95
             # Implement the category framework for the parent
            Field.__init__(self, base, category=category or QuotientFields())
        # Single-underscore method for string representation
        def _repr_(self):
100
            return "NewFrac(%s)"%repr(self.base())
101
102
        # Two methods that are implicitly used in some tests
103
104
        def base_ring(self):
            return self.base().base_ring()
105
        def characteristic(self):
106
            return self.base().characteristic()
107
108
        # Implement conversions. Do not override __call__!
109
        def _element_constructor_(self, *args, **kwds):
            if len(args)!=1:
                return self.element_class(self, *args, **kwds)
112
            x = args[0]
113
            try:
114
                 P = x.parent()
115
            except AttributeError:
116
                 return self.element_class(self, x, **kwds)
118
            if P in QuotientFields() and P != self.base():
```

```
return self.element_class(self, x.numerator(), x.denominator(), **kwds)
119
            return self.element_class(self, x, **kwds)
120
121
        # Implement coercion from the base and from fraction fields
122
123
        # over a ring that coerces into the base
        def _coerce_map_from_(self, S):
124
            if self.base().has_coerce_map_from(S):
125
                return True
126
            if S in QuotientFields():
127
                if self.base().has_coerce_map_from(S.base()):
128
                     return True
129
                if hasattr(S,'ring_of_integers') and self.base().has_coerce_map_from(S.ring_of_integers()
130
131
        # Tell how this parent was constructed, in order to enable pushout constructions
132
        def construction(self):
133
            return MyFracFunctor(), self.base()
134
135
        # return some elements of this parent
136
        def _an_element_(self):
137
            a = self.base().an_element()
138
            b = self.base_ring().an_element()
139
            if (a+b) !=0:
140
                return self(a)**2/(self(a+b)**3)
141
            if b != 0:
142
                return self(a)/self(b)**2
143
            return self(a) **2*self(b) **3
144
        def some elements(self):
145
            return [self.an_element(), self(self.base().an_element()), self(self.base_ring().an_element())
146
147
148
149
    # A construction functor for our implementation of fraction fields
   class MyFracFunctor(ConstructionFunctor):
150
        # The rank is the same for Sage's original fraction field functor
151
        rank = 5
152
        def __init__(self):
153
            # The fraction field construction is a functor
154
            # from the category of integral domains into the category of
155
            # fields
156
            # NOTE: We could actually narrow the codomain and use the
157
            # category QuotientFields()
158
            ConstructionFunctor.__init__(self, IntegralDomains(), Fields())
159
        # Applying the functor to an object. Do not override __call__!
160
        def _apply_functor(self, R):
161
            return MyFrac(R)
        # Note: To apply the functor to morphisms, implement
163
                _apply_functor_to_morphism
164
165
        # Make sure that arithmetic involving elements of Frac(R) and
166
        # MyFrac(R) works and yields elements of MyFrac(R)
167
        def merge(self, other):
            if isinstance(other, (type(self), sage.categories.pushout.FractionField)):
                return self
170
171
   # A quotient field category with additional tests.
172
   # Notes:
173
   # - Category inherits from UniqueRepresentation. Hence, there
        is only one category for given arguments.
   # - Since QuotientFieldsWithTest is a singleton (there is only
```

```
one instance of this class), we could inherit from
177
        sage.categories.category_singleton.Category_singleton
178
        rather than from sage.categories.category.Category
179
   class QuotientFieldsWithTest (Category):
180
        # Our category is a sub-category of the category of quotient fields,
181
        # by means of the following method.
182
       def super_categories(self):
183
            return [QuotientFields()]
184
185
        # Here, we could implement methods that are available for
        # all objects in this category.
        class ParentMethods:
188
            pass
189
190
        # Here, we add a new test that is available for all elements
191
        # of any object in this category.
192
        class ElementMethods:
193
            def _test_factorisation(self, **options):
194
                P = self.parent()
195
                # The methods prod() and factor() are inherited from
196
                # some other categories.
197
                assert self == P.prod([P(b)**e for b,e in self.factor()])
198
```

12.1.16 How to call a C code (or a compiled library) from Sage?

If you have some C/C++ code that you would like to call from Sage for your own use, this document is for you.

• Do you want to **contibute** to Sage by adding your interface to its code? The (more complex) instructions are available here.

Calling "hello_world()" from hello.c

Let us suppose that you have a file named ~/my_dir/hello.c containing:

```
#include <stdio.h>
void hello_world() {
    printf("Hello World\n");
}
```

In order to call this function from Sage, you must create a Cython file (i.e. a file whose extension is .pyx). Here, ~/my_dir/hello_sage.pyx contains a header describing the signature of the function that you want to call:

```
cdef extern from "hello.c":
    void hello_world()

def my_bridge_function():
    hello_world() # This is the C function from hello.c
```

You can now load this file in Sage, and call the C code though the Python function my_bridge_function:

```
sage: %runfile hello_sage.pyx
Compiling ./hello_sage.pyx...
sage: my_bridge_function()
Hello World
```

Arguments and return value

Calling function with more complex arguments and return values works the same way. To learn more about the Cython language, click here

The following example defines a function taking and returning int * pointers, and involves some memory allocation. The C code defines a function whose purpose is to return the sum of two vectors as a third vector.

```
The C file (double_vector.c)
#include <string.h>
int * sum_of_two_vectors(int n, int * vec1, int * vec2) {
  /*
     INPUT : two arrays vec1, vec2 of n integers
     OUTPUT: an array of size n equal to vec1+vec2
  int * sum = (int *) malloc(n*sizeof(int));
  int i;
  for (i=0; i<n; i++)</pre>
    sum[i] = vec1[i] + vec2[i];
  return sum;
}
The Cython file (double_vector_sage.pyx)
cdef extern from "double_vector.c":
    int * sum_of_two_vectors(int n, int * vec1, int * vec2)
from libc.stdlib cimport malloc, free
def sage_sum_of_vectors(n, list1, list2):
    cdef int * vec1 = <int *> malloc(n*sizeof(int))
    cdef int * vec2 = <int *> malloc(n*sizeof(int))
    # Fill the vectors
    for i in range(n):
        vec1[i] = list1[i]
        vec2[i] = list2[i]
    # Call the C function
    cdef int * vec3 = sum_of_two_vectors(n, vec1, vec2)
    # Save the answer in a Python object
    answer = [vec3[i] for i in range(n)]
    free (vec1)
    free (vec2)
    free (vec3)
    return answer
Call from Sage:
sage: %runfile double_vector_sage.pyx
Compiling ./double_vector_sage.pyx...
sage: sage_sum_of_vectors(3,[1,1,1],[2,3,4])
[3, 4, 5]
```

Calling code from a compiled library

The procedure is very similar again. For our purposes, we build a library from the file ~/my_dir/hello.c:

```
#include <stdio.h>
void hello_world() {
    printf("Hello World\n");
```

We also need a ~/my_dir/hello.h header file:

```
void hello_world();
```

We can now **compile it** as a library:

```
[user@localhost ~/my_dir/] gcc -c -Wall -Werror -fpic hello.c [user@localhost ~/my_dir/] gcc -shared -o libhello.so hello.o
```

The only files that we need now are hello.h and libhello.so (you can remove the others if you like). We must now indicate the location of the .so and .h files in the header of our ~/my_dir/hello_sage.pyx file:

```
#clib /home/username/my_dir/hello

cdef extern from "hello.h":
    void hello_world()

def my_bridge_function():
    hello_world() # This is the C function from hello.c
```

Note: The instruction #clib /home/username/my_dir/hello indicates that the library is actually named /home/username/my_dir/hello. Change it according to your needs. For more information about these instructions, see cython().

We can now **load** this file in Sage and **call** the function:

```
sage: %runfile hello_sage.pyx
Compiling ./hello_sage.pyx...
sage: my_bridge_function()
Hello World
```

12.1.17 Numerical Computing with Sage

This document is designed to introduce the reader to the tools in Sage that are useful for doing numerical computation. By numerical computation we essentially mean machine precision floating point computations. In particular, things such as optimization, numerical linear algebra, solving ODE's/PDE's numerically, etc. or the first part of this document the reader is only assumed to be familiar with Python/Sage. In the second section on using compiled code, the computational prerequisites increase and I assume the reader is comfortable with writing programs in C or Fortran. The third section is on mpi and parallel programming and only requires knowledge of Python, though familiarity with mpi would be helpful. Finally the last section is about 3d visualization in Sage.

In the current version of this document the reader is assumed to be familiar with the techniques of numerical analysis. The goal of this document is not to teach you numerical analysis, but to explain how to express your ideas in Sage and Python. Also this document is not meant to be comprehensive. Instead the goal is to be a road map and orient the reader to the packages relevant to numerical computation and to where they can find more information.

Numerical Tools

Sage has many different components that may be useful for numerical analysis. In particular three packages deserve mention, they are numpy, SciPy, and cvxopt. Numpy is an excellent package that provides fast array facilities to python. It includes some basic linear algebra routines, vectorized math routines, random number generators, etc. It supports a programming style similar to one would use in matlab and most matlab techniques have an analogue in numpy. SciPy builds on numpy and provides many different packages for optimization, root finding, statistics, linear algebra, interpolation, FFT and dsp tools, etc. Finally cvxopt is an optimization package which can solve linear and quadratic programming problems and also has a nice linear algebra interface. Now we will spend a bit more time on each of these packages.

Before we start let us point out http://www.scipy.org/NumPy_for_Matlab_Users, which has a comparison between matlab and numpy and gives numpy equivalents of matlab commands. If you're not familiar with matlab, thats fine, even better, it means you won't have any pre-conceived notions of how things should work. Also this http://www.scipy.org/Wiki/Documentation?action=AttachFile&do=get&target=scipy_tutorial.pdf is a very nice tutorial on SciPy and numpy which is more comprehensive than ours.

NumPy

NumPy is not imported into sage initially. To use NumPy, you first need to import it.

```
sage: import numpy
```

The basic object of computation in NumPy is an array. It is simple to create an array.

```
sage: l=numpy.array([1,2,3])
sage: l
array([1, 2, 3])
```

NumPy arrays can store any type of python object. However, for speed, numeric types are automatically converted to native hardware types (i.e., int, float, etc.) when possible. If the value or precision of a number cannot be handled by a native hardware type, then an array of Sage objects will be created. You can do calculations on these arrays, but they may be slower than using native types. When the numpy array contains Sage or python objects, then the data type is explicitly printed as object. If no data type is explicitly shown when NumPy prints the array, the type is either a hardware float or int.

The dtype attribute of an array tells you the type of the array. For fast numerical computations, you generally want this to be some sort of float. If the data type is float, then the array is stored as an array of machine floats, which takes up much less space and which can be operated on much faster.

```
sage: l=numpy.array([1.0, 2.0, 3.0])
sage: l.dtype
dtype('float64')
```

You can create an array of a specific type by specifying the dtype parameter. If you want to make sure that you are dealing with machine floats, it is good to specify dtype=float when creating an array.

```
sage: l=numpy.array([1,2,3], dtype=float)
sage: l.dtype
dtype('float64')
```

You can access elements of a NumPy array just like any list, as well as take slices

```
sage: l=numpy.array(range(10),dtype=float)
sage: 1[3]
3.0
sage: 1[3:6]
array([ 3.,  4.,  5.])
```

You can do basic arithmetic operations

```
sage: 1+1
array([ 0., 2., 4., 6., 8., 10., 12., 14., 16., 18.])
sage: 2.5*l
array([ 0., 2.5, 5., 7.5, 10., 12.5, 15., 17.5, 20., 22.5])
```

Note that 1*1 will multiply the elements of 1 componentwise. To get a dot product, use numpy.dot().

```
sage: 1*1
array([ 0., 1., 4., 9., 16., 25., 36., 49., 64., 81.])
sage: numpy.dot(1,1)
285.0
```

We can also create two dimensional arrays

This is basically equivalent to the following

The difference is that with numpy.array(), m is treated as just an array of data. In particular m*m will multiply componentwise, however with numpy.matrix(), m*m will do matrix multiplication. We can also do matrix vector multiplication, and matrix addition

If n was created with numpy.array(), then to do matrix vector multiplication, you would use numpy.dot(n, v).

All NumPy arrays have a shape attribute. This is a useful attribute to manipulate

```
sage: n = numpy.array(range(25),dtype=float)
sage: n
array([ 0., 1., 2., 3., 4., 5., 6., 7., 8., 9., 10.,
       11., 12., 13., 14., 15., 16., 17., 18., 19., 20., 21.,
       22., 23., 24.])
sage: n.shape=(5,5)
sage: n
array([[ 0.,
              1.,
                    2.,
                         3.,
                               4.],
             6.,
      [ 5.,
                   7.,
                         8.,
                               9.1,
      ſ 10.,
             11.,
                  12.,
                        13.,
                             14.1,
            16.,
                  17.,
                       18.,
                             19.],
      [ 15.,
      [ 20., 21., 22., 23., 24.]])
```

This changes the one-dimensional array into a 5×5 array.

NumPy arrays can be sliced as well

It is important to note that the sliced matrices are references to the original

```
sage: m=n[2:4,1:3]
sage: m[0,0]=100
sage: n
array([[
          0.,
                 1.,
                         2.,
                               3.,
                                       4.],
         5.,
                 6.,
                        7.,
                               8.,
                                       9.1,
      Γ
               100.,
                        12.,
       [ 10.,
                             13.,
                                      14.],
               16.,
                        17.,
                             18.,
       [ 15.,
                                      19.],
       [ 20.,
                 21.,
                        22.,
                             23.,
                                      24.]])
```

You will note that the original matrix changed. This may or may not be what you want. If you want to change the sliced matrix without changing the original you should make a copy

```
m=n[2:4,1:3].copy()
```

Some particularly useful commands are

You can see that numpy.arange() creates an array of floats increasing by 0.1 from 0 to 2. There is a useful command numpy.r_() that is best explained by example

```
sage: from numpy import r_
sage: j=numpy.complex(0,1)
sage: RealNumber=float
sage: Integer=int
sage: n=r_[0.0:5.0]
sage: n
array([ 0.,  1.,  2.,  3.,  4.])
sage: n=r_[0.0:5.0, [0.0]*5]
```

```
sage: n
array([ 0., 1., 2., 3., 4., 0., 0., 0., 0., 0.])
```

numpy.r_() provides a shorthand for constructing NumPy arrays efficiently. Note in the above 0.0:5.0 was shorthand for 0.0, 1.0, 2.0, 3.0, 4.0. Suppose we want to divide the interval from 0 to 5 into 10 intervals. We can do this as follows

```
sage: r_[0.0:5.0:11*j]
array([ 0. , 0.5, 1. , 1.5, 2. , 2.5, 3. , 3.5, 4. , 4.5, 5. ])
```

The notation 0.0:5.0:11*j expands to a list of 11 equally space points between 0 and 5 including both endpoints. Note that j is the NumPy imaginary number, but it has this special syntax for creating arrays. We can combine all of these techniques

Another useful command is numpy.meshgrid(), it produces meshed grids. As an example suppose you want to evaluate $f(x,y)=x^2+y^2$ on a an equally spaced grid with $\Delta x=\Delta y=.25$ for $0\leq x,y\leq 1$. You can do that as follows

```
sage: import numpy
sage: j=numpy.complex(0,1)
sage: def f(x,y):
\dots: return x**2+y**2
sage: from numpy import meshgrid
sage: x=numpy.r_[0.0:1.0:5*j]
sage: y=numpy.r_[0.0:1.0:5*j]
sage: xx,yy= meshgrid(x,y)
sage: xx
array([[ 0. , 0.25, 0.5 , 0.75, 1. ],
      [ 0. , 0.25, 0.5 , 0.75, 1. ],
      [ 0. , 0.25, 0.5 , 0.75, 1. ],
      [ 0. , 0.25, 0.5 , 0.75, 1. ],
      [ 0. , 0.25, 0.5 , 0.75, 1.
                                    11)
sage: yy
array([[ 0. , 0. ,
                    0.,
                          0. , 0. ],
      [0.25, 0.25, 0.25, 0.25, 0.25],
             0.5 , 0.5 ,
      [ 0.5 ,
                          0.5 ,
                                0.5 ],
      [ 0.75, 0.75, 0.75, 0.75, 0.75],
      [1., 1., 1., 1., 1.]])
sage: f(xx,yy)
array([[ 0. , 0.0625, 0.25 , 0.5625, 1.
      [ 0.0625, 0.125 , 0.3125, 0.625 , 1.0625],
      [ 0.25 , 0.3125, 0.5 , 0.8125, 1.25 ],
      [0.5625, 0.625, 0.8125, 1.125, 1.5625],
      [ 1. , 1.0625, 1.25 , 1.5625,
                                       2.
                                            11)
```

You can see that numpy.meshgrid() produces a pair of matrices, here denoted xx and yy, such that (xx[i,j],yy[i,j]) has coordinates (x[i],y[j]). This is useful because to evaluate f over a grid, we only need to evaluate it on each pair of entries in xx, yy. Since NumPy automatically performs arithmetic operations on arrays componentwise, it is very easy to evaluate functions over a grid with very little code.

A useful module is the numpy.linalg module. If you want to solve an equation Ax = b do

```
sage: import numpy
sage: from numpy import linalg
sage: A=numpy.random.randn(5,5)
sage: b=numpy.array(range(1,6))
sage: x=linalg.solve(A,b)
sage: numpy.dot(A,x)
array([ 1., 2., 3., 4., 5.])
```

This creates a random 5x5 matrix A, and solves Ax = b where b = [0.0, 1.0, 2.0, 3.0, 4.0]. There are many other routines in the numpy.linalg module that are mostly self-explanatory. For example there are qr and lu routines for doing QR and LU decompositions. There is also a command eigs for computing eigenvalues of a matrix. You can always do <function name>? to get the documentation which is quite good for these routines.

Hopefully this gives you a sense of what NumPy is like. You should explore the package as there is quite a bit more functionality.

SciPy

Again I recommend this http://www.scipy.org/Wiki/Documentation?action=AttachFile&do=get&target=scipy_tutorial.pdf. There are many useful SciPy modules, in particular scipy.optimize, scipy.stats, scipy.linalg, scipy.linsolve, scipy.sparse, scipy.integrate, scipy.ftpack, scipy.signal, scipy.special. Most of these have relatively good documentation and often you can figure out what things do from the names of functions. I recommend exploring them. For example if you do

```
sage: import scipy
sage: from scipy import optimize
Then
sage: optimize.[tab]
```

will show a list of available functions. You should see a bunch of routines for finding minimum of functions. In particular if you do

```
sage: optimize.fmin_cg?
```

you find it is a routine that uses the conjugate gradient algorithm to find the minima of a function.

```
sage: scipy.special.[tab]
```

will show all the special functions that SciPy has. Spending a little bit of time looking around is a good way to familiarize yourself with SciPy. One thing that is sort of annoying, is that often if you do scipy.:math: \langle tab \rangle . You won't see a module that is importable. For example scipy.:math: \langle tab \rangle will not show a signal module but

```
sage: from scipy import signal
and then
signal.[tab]
```

will show you a large number of functions for signal processing and filter design. All the modules I listed above can be imported even if you can't see them initially.

scipy.integrate This module has routines related to numerically solving ODE's and numerical integration. Lets give an example of using an ODE solver. Suppose you want to solve the ode

$$x''(t) + ux'(t)(x(t)^{2} - 1) + x(t) = 0$$

which as a system reads

```
x' = y
y' = -x + \mu y(1 - x^2).
```

The module we want to use is odeint in scipy.integrate. We can solve this ode, computing the value of (y, y'), at 1000 points between 0, and 100 using the following code.

We could plot the solution if we wanted by doing

```
sage: pts = [(x[i],y[i][0]) for i in range(len(x))]
sage: point2d(pts).show()
```

Optimization The Optimization module has routines related to finding roots, least squares fitting, and minimization. \(\triangle \text{To be Written } \)

Cvxopt

Cvxopt provides many routines for solving convex optimization problems such as linear and quadratic programming packages. It also has a very nice sparse matrix library that provides an interface to umfpack (the same sparse matrix solver that matlab uses), it also has a nice interface to lapack. For more details on cvxopt please refer to its documentation at http://cvxopt.org/userguide/index.html

Sparse matrices are represented in triplet notation that is as a list of nonzero values, row indices and column indices. This is internally converted to compressed sparse column format. So for example we would enter the matrix

$$\left(\begin{array}{cccccc}
2 & 3 & 0 & 0 & 0 \\
3 & 0 & 4 & 0 & 6 \\
0 & -1 & -3 & 2 & 0 \\
0 & 0 & 1 & 0 & 0 \\
0 & 4 & 2 & 0 & 1
\end{array}\right)$$

by

```
sage: import numpy
sage: from cvxopt.base import spmatrix
sage: from cvxopt.base import matrix as m
sage: from cvxopt import umfpack
sage: Integer=int
sage: V = [2,3, 3,-1,4, 4,-3,1,2, 2, 6,1]
sage: I = [0,1, 0, 2,4, 1, 2,3,4, 2, 1,4]
sage: J = [0,0, 1, 1,1, 2, 2,2,2, 3, 4,4]
sage: A = spmatrix(V,I,J)
```

To solve an equation AX = B, with B = [1, 1, 1, 1, 1], we could do the following.

```
sage: B = numpy.array([1.0]*5)
sage: B.shape=(5,1)
sage: print(B)
[[1.]
[ 1.]
[ 1.]
[ 1.]
[ 1.]]
sage: print(A)
                          0
[ 2.00e+00 3.00e+00 0
                                     0 ]
6.00e+00]
   0 -1.00e+00 -3.00e+00 2.00e+00
                                   0 ]
          0 1.00e+00 0
                                      0
                                          1
Γ
    0
         4.00e+00 2.00e+00
                           0
                                   1.00e+001
[
sage: C=m(B)
sage: umfpack.linsolve(A,C)
sage: print(C)
[ 5.79e-01]
[-5.26e-02]
[ 1.00e+00]
[ 1.97e+00]
[-7.89e-01]
```

Note the solution is stored in B afterward. also note the m(B), this turns our numpy array into a format cvxopt understands. You can directly create a cvxopt matrix using cvxopt's own matrix command, but I personally find numpy arrays nicer. Also note we explicitly set the shape of the numpy array to make it clear it was a column vector.

We could compute the approximate minimum degree ordering by doing

```
sage: RealNumber=float
sage: Integer=int
sage: from cvxopt.base import spmatrix
sage: from cvxopt import amd
sage: A=spmatrix([10,3,5,-2,5,2],[0,2,1,2,2,3],[0,0,1,1,2,3])
sage: P=amd.order(A)
sage: print(P)
[ 1]
[ 0]
[ 2]
[ 3]
```

For a simple linear programming example, if we want to solve

```
\begin{array}{ll} \text{minimze} & -4x_1 - 5x_2 \\ \text{subject to} & 2x_1 + x_2 \leq 3 \\ & x_1 + 2x_2 \leq 3 \\ & x_1 \geq 0 \\ & x_2 \geq 0 \end{array}
```

```
sage: RealNumber=float
sage: Integer=int
sage: from cvxopt.base import matrix as m
sage: from cvxopt import solvers
sage: c = m([-4., -5.])
sage: G = m([[2., 1., -1., 0.], [1., 2., 0., -1.]])
sage: h = m([3., 3., 0., 0.])
sage: sol = solvers.lp(c,G,h) #random
```

```
pcost
                                  pres
                                                k/t
               dcost.
                                         dres
                           gap
0: -8.1000e+00 -1.8300e+01 4e+00 0e+00 8e-01 1e+00
1: -8.8055e+00 -9.4357e+00 2e-01 1e-16 4e-02
                                                3e - 02
2: -8.9981e+00 -9.0049e+00 2e-03
                                  1e-16
                                         5e-04
                                                4e - 04
3: -9.0000e+00 -9.0000e+00 2e-05
                                  3e-16
                                         5e-06
4: -9.0000e+00 -9.0000e+00 2e-07 1e-16
                                         5e-08
sage: print sol['x'] # ... below since can get -00 or +00 depending on architecture
[ 1.00e...00]
[ 1.00e+00]
```

Using Compiled Code Interactively

This section is about using compiled code in Sage. However, since Sage is built on top of Python most of this is valid for Python in general. The exception is that these notes assume you are using Sage's interface to f2py which makes it more convenient to work with f2py interactively. You should look at the f2py website for information on using the command line f2py tool. The ctypes and weave example will work in any recent Python install (weave is not part of Python so you will have to install it separately). If you are using Sage then weave, ctypes, and f2py are all there already.

Firstly why would we want to write compiled code? Obviously, because its fast, far faster than interpreted Python code. Sage has very powerful facilities that allow one to interactively call compiled code written in C or Fortran. In fact there 2-4 ways to do this depending on exactly what you want to accomplish. One way is to use Cython. Cython is a language that is a hybrid of C and Python based on Pyrex. It has the ability to call external shared object libraries and is very useful for writing Python extension modules. Cython/Pyrex is covered in detail elsewhere in the Sage documentation.

Suppose that you really want to just write Python code, but there is some particularly time intensive piece of your code that you would like to either write in C/Fortran or simply call an external shared library to accomplish. In this case you have three options with varying strengths and weaknesses.

Note that before you try to use compiled code to speed up your bottleneck make sure there isn't an easier way. In particular, first try to vectorize, that is express your algorithm as arithmetic on vectors or numpy arrays. These arithmetic operations are done directly in C so will be very fast. If your problem does not lend itself to being expressed in a vectorized form them read on.

Before we start let us note that this is in no way a complete introduction to any of the programs we discuss. This is more meant to orient you to what is possible and what the different options will feel like.

f2py

F2py is a very nice package that automatically wraps fortran code and makes it callable from Python. The Fibonacci examples are taken from the f2py webpage http://cens.ioc.ee/projects/f2py2e/.

From the notebook the magic %fortran will automatically compile any fortran code in a cell and all the subroutines will become callable functions (though the names will be converted to lowercase.) As an example paste the following into a cell. It is important that the spacing is correct as by default the code is treated as fixed format fortran and the compiler will complain if things are not in the correct column. To avoid this, you can write fortran 90 code instead by making your first line !f90. There will be an example of this later.

```
C
INTEGER N
REAL*8 A(N)
DO I=1,N
IF (I.EQ.1) THEN
A(I) = 0.0D0
ELSEIF (I.EQ.2) THEN
A(I) = 1.0D0
ELSE
A(I) = A(I-1) + A(I-2)
ENDIF
ENDDO
END
C END FILE FIB1.F
```

Now evaluate it. It will be automatically compiled and imported into Sage (though the name of imported function will be lowercase). Now we want to try to call it, we need to somehow pass it an array A, and the length of the array N. The way it works is that numpy arrays will be automatically converted to fortran arrays, and Python scalars converted to fortran scalars. So to call fib we do the following.

```
import numpy
m=numpy.array([0]*10,dtype=float)
print(m)
fib(m,10)
print(m)
```

Note that fortran is a function that can be called on any string. So if you have a fortran program in a file my prog.f. Then you could do the following

```
f=open('my_prog.f','r')
s=f.read()
fortran(s)
```

Now all the functions in my prog.f are callable.

It is possible to call external libraries in your fortran code. You simply need to tell f2py to link them in. For example suppose we wish to write a program to solve a linear equation using lapack (a linear algebra library). The function we want to use is called dgesv and it has the following signature.

```
SUBROUTINE DGESV( N, NRHS, A, LDA, IPIV, B, LDB, INFO )
           (input) INTEGER
  Ν
           The number of linear equations, i.e., the order of the
           matrix A. N >= 0.
  NRHS
           (input) INTEGER
           The number of right hand sides, i.e., the number of columns
           of the matrix B. NRHS >= 0.
           (input/output) DOUBLE PRECISION array, dimension (LDA, N)
  Α
           On entry, the N-by-N coefficient matrix A.
           On exit, the factors L and U from the factorization
           A = P*L*U; the unit diagonal elements of L are not stored.
           (input) INTEGER
  LDA
           The leading dimension of the array A. LDA >= \max(1, N).
```

```
TPTV
        (output) INTEGER array, dimension (N)
        The pivot indices that define the permutation matrix P;
        row i of the matrix was interchanged with row IPIV(i).
В
        (input/output) DOUBLE PRECISION array, dimension (LDB, NRHS)
        On entry, the N-by-NRHS matrix of right hand side matrix B.
        On exit, if INFO = 0, the N-by-NRHS solution matrix X.
        (input) INTEGER
LDB
        The leading dimension of the array B. LDB \geq \max(1, N).
        (output) INTEGER
        = 0: successful exit
        < 0: if INFO = -i, the i-th argument had an illegal value
        > 0: if INFO = i, U(i,i) is exactly zero. The factorization
              has been completed, but the factor U is exactly
              singular, so the solution could not be computed.
```

we could do the following. Note that the order that library are in the list actually matters as it is the order in which they are passed to gcc. Also fortran.libraries is simply a list of names of libraries that are linked in. You can just directly set this list. So that fortran.libraries=['lapack','blas']is equivalent to the following.

```
fortran.add_library('lapack')
fortran.add_library('blas')
```

Now

```
%fortran
!f90
Subroutine LinearEquations(A,b,n)
Integer n
Real*8 A(n,n), b(n)
Integer i, j, pivot(n), ok
call DGESV(n, 1, A, n, pivot, b, n, ok)
end
```

There are a couple things to note about this. As we remarked earlier, if the first line of the code is !f90, then it will be treated as fortran 90 code and does not need to be in fixed format. To use the above try

```
a=numpy.random.randn(10,10)
b=numpy.array(range(10),dtype=float)
x=b.copy()
linearequations(a,x,10)
numpy.dot(a,x)
```

This will solve the linear system ax=b and store the result in b. If your library is not in Sage's local/lib or in your path you can add it to the search path using

```
fortran.add_library_path('path').
```

You can also directly set fortran.library paths by assignment. It should be a list of paths (strings) to be passed to gcc. To give you an idea of some more things you can do with f2py, note that using intent statements you can control the way the resulting Python function behaves a bit bitter. For example consider the following modification of our original fibonacci code.

```
C FILE: FIB3.F
SUBROUTINE FIB(A,N)
```

```
C
С
      CALCULATE FIRST N FIBONACCI NUMBERS
С
      INTEGER N
      REAL * 8 A(N)
Cf2py intent(in) n
Cf2py intent(out) a
Cf2py depend(n) a
      DO I=1, N
         IF (I.EQ.1) THEN
            A(I) = 0.0D0
         ELSEIF (I.EQ.2) THEN
            A(I) = 1.0D0
         ELSE
            A(I) = A(I-1) + A(I-2)
         ENDIF
      ENDDO
      END
C END FILE FIB3.F
```

Note the comments with the intent statements. This tells f2py that n is an input parameter and a is the output. This is called as

```
a=fib(10)
```

In general you will pass everything declared intent(in) to the fortran function and everything declared intent(out) will be returned in a tuple. Note that declaring something intent(in) means you only care about its value before the function is called not afterwards. So in the above n tells us how many fiboncci numbers to compute we need to specify this as an input, however we don't need to get n back as it doesn't contain anything new. Similarly A is intent(out) so we don't need A to have an specific value beforehand, we just care about the contents afterwards. F2py generates a Python function so you only pass those declared intent(in) and supplies empty workspaces for the remaining arguments and it only returns those that are intent(out). All arguments are intent(in) by default.

Consider now the following

```
%fortran
    Subroutine Rescale(a,b,n)
    Implicit none
    Integer n,i,j
    Real*8 a(n,n), b
    do i = 1,n
        do j=1,n
        a(i,j)=b*a(i,j)
    end do
end
end
```

You might be expecting Rescale(a,n) to rescale a numpy matrix a. Alas this doesn't work. Anything you pass in is unchanged afterwards. Note that in the fibonacci example above, the one dimensional array was changed by the fortran code, similarly the one dimensional vector b was replaced by its solution in the example where we called lapack while the matrix A was not changed even then dgesv says it modifies the input matrix. Why does this not happen with the two dimensional array. Understanding this requires that you are aware of the difference between how fortran and C store arrays. Fortran stores a matrices using column ordering while C stores them using row ordering. That is the matrix

$$\left(\begin{array}{ccc} 0 & 1 & 2 \\ 3 & 4 & 5 \end{array}\right)$$

is stored as

```
(012345) in C (031425) in Fortran
```

One dimensional arrays are stored the same in C and Fortran. Because of this f2py allows the fortran code to operate on one dimensional vectors in place, so your fortran code will change one dimensional numpy arrays passed to it. However, since two dimensional arrays are different by default f2py copies the numpy array (which is stored in C format) into a second array that is in the fortran format (i.e. takes the transpose) and that is what is passed to the fortran function. We will see a way to get around this copying later. First let us point one way of writing the rescale function.

%fortran

```
Subroutine Rescale(a,b,n)
Implicit none
Integer n,i,j
Real*8 a(n,n), b

Cf2py intent(in,out) a
do i = 1,n
do j=1,n
a(i,j)=b*a(i,j)
end do
end do
end
```

Note that to call this you would use

```
b=rescale(a, 2.0).
```

Note here I am not passing in n which is the dimension of a. Often f2py can figure this out. This is a good time to mention that f2py automatically generates some documentation for the Python version of the function so you can check what you need to pass to it and what it will return. To use this try

```
rescale?
```

The intent(in,out) directives tells f2py to take the contents of a at the end of the subroutine and return them in a numpy array. This still may not be what you want. The original a that you pass in is unmodified. If you want to modify the original a that you passed in use intent(inout). This essentially lets your fortran code work with the data inplace.

%fort.ran

```
Subroutine Rescale(a,b,n)
Implicit none
Integer n,i,j
Real*8 a(n,n), b

Cf2py intent(inout) a
do i = 1,n
do j=1,n
a(i,j)=b*a(i,j)
end do
end do
end
```

If you wish to have fortran code work with numpy arrays in place, you should make sure that your numpy arrays are stored in fortran's format. You can ensure this by using the order='FORTRAN' keyword when creating the arrays, as follows.

```
a=numpy.array([[1,2],[3,4]],dtype=float,order='FORTRAN')
rescale(a,2.0)
```

After this executes, a will have the rescaled version of itself. There is one final version which combines the previous two.

%fortran

```
Subroutine Rescale(a,b,n)
Implicit none
Integer n,i,j
Real*8 a(n,n), b

Cf2py intent(in,out,overwrite) a
do i = 1,n
do j=1,n
a(i,j)=b*a(i,j)
end do
end do
end
```

The (in,out,overwite) intent says that if *a* is in FORTRAN ordering we work in place, however if its not we copy it and return the contents afterwards. This is sort of the best of both worlds. Note that if you are repeatedly passing large numpy arrays to fortran code, it is very important to avoiding copying the array by using (inout) or (in,out,overwrite). Remember though that your numpy array must use Fortran ordering to avoid the copying.

For more examples and more advanced usage of F2py you should refer to the f2py webpage http://cens.ioc.ee/projects/f2py2e/. The command line f2py tool which is referred to in the f2py documentation can be called from the Sage shell using

!f2py

More Interesting Examples with f2py

Let us now look at some more interesting examples using f2py. We will implement a simple iterative method for solving laplace's equation in a square. Actually, this implementation is taken from http://www.scipy.org/PerformancePython?highlight=%28performance%29 page on the scipy website. It has lots of information on implementing numerical algorithms in python.

The following fortran code implements a single iteration of a relaxation method for solving Laplace's equation in a square.

```
%fortran
```

```
subroutine timestep(u,n,m,dx,dy,error)
double precision u(n,m)
double precision dx,dy,dx2,dy2,dnr_inv,tmp,diff
integer n,m,i,j
cf2py intent(in) :: dx,dy
cf2py intent(in,out) :: u
cf2py intent(out) :: error
cf2py intent(hide) :: n,m
dx2 = dx*dx
dy2 = dy*dy
dnr_inv = 0.5d0 / (dx2+dy2)
error = 0d0
do 200,j=2,m-1
do 100,i=2,n-1
```

If you do

timestep?

You find that you need pass timestep a numpy array u, and the grid spacing dx,dy and it will return the updated u, together with an error estimate. To apply this to actually solve a problem use this driver code

```
import numpy
j=numpy.complex(0,1)
num_points=50
u=numpy.zeros((num_points,num_points),dtype=float)
pi_c=float(pi)
x=numpy.r_[0.0:pi_c:num_points*j]
u[0,:]=numpy.sin(x)
u[num_points-1,:]=numpy.sin(x)
def solve_laplace(u,dx,dy):
    iter =0
    err = 2
    while(iter <10000 and err>le-6):
        (u,err)=timestep(u,dx,dy)
        iter+=1
    return (u,err,iter)
```

Now call the routine using

```
(sol,err,iter)=solve_laplace(u,pi_c/(num_points-1),pi_c/(num_points-1))
```

This solves the equation with boundary conditions that the right and left edge of the square are half an oscillation of the sine function. With a 51x51 grid that we are using I find that it takes around .2 s to solve this requiring 2750 iterations. If you have the gnuplot package installed (use optional packages() to find its name and install package to install it), then you can plot this using

import Gnuplot

```
g=Gnuplot.Gnuplot(persist=1)
g('set parametric')
g('set data style lines')
g('set hidden')
g('set contour base')
g('set zrange [-.2:1.2]')
data=Gnuplot.GridData(sol,x,x,binary=0)
g.splot(data)
```

To see what we have gained by using f2py let us compare the same algorithm in pure python and a vectorized version using numpy arrays.

```
def slowTimeStep(u,dx,dy):
    """Takes a time step using straight forward Python loops."""
    nx, ny = u.shape
```

```
dx2, dy2 = dx**2, dy**2
    dnr_inv = 0.5/(dx2 + dy2)
    err = 0.0
    for i in range (1, nx-1):
        for j in range(1, ny-1):
            tmp = u[i,j]
            u[i,j] = ((u[i-1, j] + u[i+1, j])*dy2 +
                      (u[i, j-1] + u[i, j+1])*dx2)*dnr_inv
            diff = u[i,j] - tmp
            err += diff*diff
    return u, numpy.sqrt(err)
def numpyTimeStep(u,dx,dy):
    dx2, dy2 = dx**2, dy**2
    dnr_inv = 0.5/(dx2 + dy2)
    u_old=u.copy()
    # The actual iteration
    u[1:-1, 1:-1] = ((u[0:-2, 1:-1] + u[2:, 1:-1])*dy2 +
                     (u[1:-1,0:-2] + u[1:-1, 2:])*dx2)*dnr_inv
    v = (u - u_old).flat
    return u, numpy.sqrt(numpy.dot(v, v))
```

You can try these out by changing the timestep function used in our driver routine. The python version is slow even on a 50x50 grid. It takes 70 seconds to solve the system in 3000 iterations. It takes the numpy routine 2 seconds to reach the error tolerance in around 5000 iterations. In contrast it takes the f2py routine around .2 seconds to reach the error tolerance using 3000 iterations. I should point out that the numpy routine is not quite the same algorithm since it is a jacobi iteration while the f2py one is gauss-seidel. This is why the numpy version requires more iterations. Even accounting for this you can see the f2py version appears to be around 5 times faster than the numpy version. Actually if you try this on a 500x500 grid I find that it takes the numpy routine 30 seconds to do 500 iterations while it only takes about 2 seconds for the f2py to do this. So the f2py version is really about 15 times faster. On smaller grids each actual iteration is relatively cheap and so the overhead of calling f2py is more evident, on larger examples where the iteration is expensive, the advantage of f2py is clear. Even on the small example it is still very fast. Note that a 500x500 grid in python would take around half an hour to do 500 iterations.

To my knowledge the fastest that you could implement this algorithm in matlab would be to vectorize it exactly like the numpy routine we have. Vector addition in matlab and numpy are comparable. So unless there is some trick I don't know about, using f2py you can interactively write code 15 times faster than anything you could write in matlab (Please correct me if I'm wrong). You can actually make the f2py version a little bit faster by using intent(in,out,overwrite) and creating the initial numpy array using order='FORTRAN'. This eliminates the unnecessary copying that is occurring in memory.

Weave

Weave is a tool that does for C/C++ what f2py does for fortran (though we should note it is also possible to wrap C code using f2py). Suppose we have some data stored in numpy arrays and we want to write some C/C++ code to do something with that data that needs to be fast. For a trivial example, let us write a function that sums the contents of a numpy array

```
sage: from scipy import weave
sage: from scipy.weave import converters
```

```
def my_sum(a):
    n=int(len(a))
    code="""
    int i;
    long int counter;
    counter =0;
    for(i=0;i<n;i++)
    {
        counter=counter+a(i);
    }
    return_val=counter;
    """

    err=weave.inline(code,['a','n'],type_converters=converters.blitz,compiler='gcc')
    return err</pre>
```

To call this function do

```
import numpy
a = numpy.array(range(60000))
time my_sum(a)
time sum(range(60000))
```

The first time the weave code executes the code is compiled, from then on, the execution is immediate. You should find that python's built-in sum function is comparable in speed to what we just wrote. Let us explain some things about this example. As you can see, to use weave you create a string containing pure C/C++ code. Then you call weave inline on it. You pass to weave the string with the code, as well as a list of python object that it is to automatically convert to C variables. So in our case we can refer to the python objects a and n inside of weave. Numpy arrays are accessed by a(i) if they are one-dimensional or a(i,j) if they are two dimensional. Of course we cannot use just any python object, currently weave knows about all python numerical data types such as ints and floats, as well as numpy arrays. Note that numpy arrays do not become pointers in the C code (which is why they are accessed by () and not []. If you need a pointer you should copy the data into a pointer. Next is a more complicated example that calls lapack to solve a linear system ax=b.

```
def weave_solve(a,b):
    n = len(a[0])
    x = numpy.array([0]*n,dtype=float)
    support_code="""
    #include <stdio.h>
    extern "C" {
    void dgesv_(int *size, int *flag,double* data,int*size,
                int*perm, double*vec, int*size, int*ok);
    . . . .
    code="""
        int i, j;
        double* a c;
        double* b_c;
        int size;
        int flag;
        int* p;
        int ok;
        size=n;
        flag=1;
        a_c= (double *)malloc(sizeof(double)*n*n);
```

```
b_c= (double *)malloc(sizeof(double)*n);
    p = (int*)malloc(sizeof(int)*n);
    for(i=0;i<n;i++)
       b_c[i]=b(i);
       for (j=0; j<n; j++)
         a_c[i*n+j]=a(i,j);
    dgesv_(&size, &flag, a_c, &size, p, b_c, &size, &ok);
    for(i=0;i<n;i++)
       x(i)=b_c[i];
    free(a_c);
    free (b_c);
    free(p);
0.00
libs=['lapack','blas','g2c']
dirs=['/media/sdb1/sage-2.6.linux32bit-i686-Linux']
vars = ['a', 'b', 'x', 'n']
weave.inline(code, vars, support_code=support_code, libraries=libs, library_dirs=dirs, \)
type_converters=converters.blitz,compiler='gcc')
return x
```

Note that we have used the support_code argument which is additional C code you can use to include headers and declare functions. Note that inline also can take all distutils compiler options which we used here to link in lapack.

```
def weaveTimeStep(u, dx, dy):
    """Takes a time step using inlined C code -- this version uses
    blitz arrays."""
    nx, ny = u.shape
    dx2, dy2 = dx * *2, dy * *2
    dnr_inv = 0.5/(dx2 + dy2)
    code = """
           double tmp, err, diff,dnr_inv_;
           dnr_inv_=dnr_inv;
           err = 0.0;
           for (int i=1; i<nx-1; ++i) {
               for (int j=1; j< ny-1; ++j) {
                   tmp = u(i,j);
                   u(i,j) = ((u(i-1,j) + u(i+1,j))*dy2 +
                              (u(i,j-1) + u(i,j+1))*dx2)*dnr_inv_;
                   diff = u(i,j) - tmp;
                   err += diff*diff;
           }
           return_val = sqrt(err);
    # compiler keyword only needed on windows with MSVC installed
    err = weave.inline(code, ['u', 'dx2', 'dy2', 'dnr_inv', 'nx','ny'],
                       type_converters = converters.blitz,
                       compiler = 'gcc')
    return u,err
```

Using our previous driver you should find that this version takes about the same amount of time as the f2py version around .2 seconds to do 2750 iterations.

For more about weave see http://www.scipy.org/Weave

Ctypes

Ctypes is a very interesting python package which lets you import shared object libraries into python and call them directly. I should say that even though this is called ctypes, it can be used just as well to call functions from libraries written in fortran. The only complication is you need to know what a fortran function looks like to C. This is simple everything is a pointer, so if your fortran function would be called as foo(A,N) where A is an array and N is its length, then to call it from C it takes a pointer to an array of doubles and a pointer to an int. The other thing to be aware of is that from C, fortran functions usually have an underscore appended. That is, a fortran function foo would appear as foo from C (this is usually the case but is compiler dependent). Having said this, the following examples are in C.

As an example suppose you write the following simple C program

```
#include <stdio.h>
int sum(double *x,int n)
{
   int i;
   double counter;
   counter = 0;
   for(i=0;i<n;i++)
      {
      counter=counter+x[i];
    }
   return counter;
}</pre>
```

which you want to call from python. First make a shared object library by doing (at the command line)

```
gcc -c sum.c
gcc -shared -o sum.so sum.o
```

Note that on OSX -shared should be replaced by -dynamiclib and sum.so should be called sum.dylib Then you can do

```
from ctypes import *
my_sum=CDLL('sum.so')
a=numpy.array(range(10),dtype=float)
my_sum.sum(a.ctypes.data_as(c_void_p),int(10))
```

Note here that a.ctypes.data as(c void p) returns a ctypes object that is void pointer to the underlying array of a. Note that even though sum takes a double*, as long as we have a pointer to the correct data it doesn't matter what its type is since it will be automatically cast.

Note that actually there are other ways to pass in the required array of doubles. For example

```
a=(c_double*10)()
for i in range(10):
    a[i]=i
my_sum.sum(a,int(10))
```

This example only uses ctypes. Ctypes has wrappers for C data types so for example

```
a=c\_double(10.4)
```

would create a ctypes double object which could be passed to a C function. Note that there is a byref function that lets you pass parameters by reference. This is used in the next example. c double*10, is a python object that represents an array of 10 doubles and

```
a = (c_double * 10) ()
```

sets a equal to an array of 10 doubles. I find this method is usually less useful than using numpy arrays when the data is mathematical as numpy arrays are more well integrated into python and sage.

Here is an example of using ctypes to directly call lapack. Note that this will only work if you have a lapack shared object library on your system. Also on linux the file would be liblapack.so and you will probably use dgesv (OSX use CLAPACK hence the lack of the underscore).

For completeness, let us consider a way to solve the laplace equation using C types. Suppose you have written a simple solver in C and you want to call it from python so you can easily test different boundary conditions. Your C program might look like this.

```
#include <math.h>
#include <stdio.h>
double timestep(double *u,int nx,int ny,double dx,double dy)
  double tmp, err, diff, dx2, dy2, dnr_inv;
  dx2=dx*dx;
  dy2=dy*dy;
  dnr_inv=0.5/(dx2+dy2);
  err = 0.0;
  int i, j;
for (i=1; i<nx-1; ++i) {</pre>
  for (j=1; j<ny-1; ++j) {</pre>
    tmp = u[i*nx+j];
    u[i*nx+j] = ((u[(i-1)*nx+j] + u[(i+1)*nx+j])*dy2 +
          (u[i*nx+j-1] + u[i*nx+j+1])*dx2)*dnr_inv;
    diff = u[i*nx+j] - tmp;
    err += diff*diff;
  }
}
return sqrt(err);
double solve_in_C(double *u,int nx,int ny,double dx,double dy)
  double err;
  int iter;
  iter = 0;
  err = 1;
    while (iter <10000 && err > 1e-6)
    err=timestep(u,nx,ny,dx,dy);
```

```
iter++;
}
return err;
}
```

We can compile it by running at the command line

```
gcc -c laplace.c
gcc -shared -o laplace.so laplace.o
```

Now in sage (notebook or command line) execute

```
from ctypes import *
laplace=CDLL('/home/jkantor/laplace.so')
laplace.timestep.restype=c_double
laplace.solve_in_C.restype=c_double
import numpy
u=numpy.zeros((51,51),dtype=float)
pi_c=float(pi)
x=numpy.arange(0,pi_c+pi_c/50,pi_c/50,dtype=float)
u[0,:]=numpy.sin(x)
u[50,:]=numpy.sin(x)
def solve(u):
 iter = 0
  err = 2
  n=c_int(int(51))
  pi_c=float(pi/50)
  dx=c_double(pi_c)
  while(iter <5000 and err>1e-6):
     err=laplace.timestep(u.ctypes.data_as(c_void_p),n,n,dx,dx)
     if(iter %50==0):
       print((err,iter))
  return (u,err,iter)
```

Note the line laplace.timestep.restype=c double. By default ctypes assumes the return values are ints. If they are not you need to tell it by setting restype to the correct return type. If you execute the above code, then solve(u) will solve the system. It is comparable to the weave or fortran solutions taking around .2 seconds. Alternatively you could do

```
n=c_int(int(51))
dx=c_double(float(pi/50))
laplace.solve_in_C(n.ctypes.data_as(c_void_p),n,n,dx,dx)
```

which computes the solution entirely in C. This is very fast. Admittedly we could have had our fortran or weave routines do the entire solution at the C/Fortran level and we would have the same speed.

As I said earlier you can just as easily call a shared object library that is written in Fortran using ctypes. The key point is it must be a shared object library and all fortran arguments are passed by reference, that is as pointers or using byref. Also even though we used very simple data types, it is possible to deal with more complicated C structures. For this and more about ctypes see http://python.net/crew/theller/ctypes/

More complicated ctypes example

Here we will look at a more complicated example. First consider the following C code.

```
#include <stdio.h>
#include <stdlib.h>
struct double_row_element_t {
  double value;
  int col_index;
  struct double_row_element_t * next_element;
typedef struct double_row_element_t double_row_element;
typedef struct {
 int nrows;
  int ncols;
 int nnz;
  double_row_element** rows;
} double_sparse_matrix;
double_sparse_matrix * initialize_matrix(int nrows, int ncols)
  int i;
  double_sparse_matrix* new_matrix;
  new_matrix = (double_sparse_matrix *) malloc(sizeof(double_sparse_matrix));
  new_matrix->rows= (double_row_element **) malloc(sizeof(double_row_element *)*nrows);
  for (i=0; i<nrows; i++)</pre>
   {
      (new_matrix->rows)[i]=(double_row_element *) malloc(sizeof(double_row_element));
      (new_matrix->rows)[i]->value=0;
      (new_matrix->rows)[i]->col_index=0;
      (new_matrix->rows)[i]->next_element = 0;
  new_matrix->nrows=nrows;
  new_matrix->ncols=ncols;
  new_matrix->nnz=0;
  return new_matrix;
int free_matrix(double_sparse_matrix * matrix)
  int i;
  double_row_element * next_element;
  double_row_element* current_element;
  for (i=0; i < matrix -> nrows; i++)
    {
      current_element = (matrix->rows)[i];
      while (current_element->next_element!=0)
      next_element=current_element->next_element;
      free(current_element);
      current_element=next_element;
      free(current_element);
  free (matrix->rows);
```

```
free (matrix);
 return 1;
int set_value(double_sparse_matrix * matrix,int row, int col, double value)
 int i;
  i=0;
  double_row_element* current_element;
  double_row_element* new_element;
  if(row> matrix->nrows || col > matrix->ncols || row <0 || col <0)</pre>
   return 1:
  current_element = (matrix->rows)[row];
  while(1)
    {
      if (current_element->col_index==col)
    current_element->value=value;
    return 0;
    }
      else
    if (current_element->next_element!=0)
          if(current_element->next_element->col_index <=col)</pre>
        current_element = current_element->next_element;
      else
        if(current_element->next_element->col_index > col)
        new_element = (double_row_element *) malloc(sizeof(double_row_element));
        new_element->value=value;
        new_element->col_index=col;
        new_element->next_element=current_element->next_element;
        current_element->next_element=new_element;
        return 0;
    }
    else
        new_element = (double_row_element *) malloc(sizeof(double_row_element));
        new_element->value=value;
        new_element->col_index=col;
        new_element->next_element=0;
        current_element->next_element=new_element;
        break;
    }
  return 0;
double get_value(double_sparse_matrix* matrix, int row, int col)
```

```
double_row_element * current_element;
     if(row> matrix->nrows || col > matrix->ncols || row <0 || col <0)</pre>
          return 0.0;
     current_element = (matrix->rows)[row];
     while (1)
               if (current_element->col_index==col)
               return current_element->value;
               else
           {
                if(current_element->col_index<col && current_element->next_element !=0)
                    current_element=current_element->next_element;
               else
                     if(current_element->col_index >col || current_element ->next_element==0)
                             return 0;
           }
}
Put it in a file called linked_list_sparse.c and compile it using
gcc -c linked_list_sparse.c
gcc -shared -o linked_list_sparse.so linked_list_sparse.o
Next consider the following python helper code.
from ctypes import *
class double_row_element (Structure):
          pass
double_row_element._fields_=[("value",c_double),("col_index",c_int),("next_element",POINTER(double_row_element",POINTER(double_row_element",POINTER(double_row_element),("col_index",c_int),("next_element",POINTER(double_row_element),("col_index",c_int),("next_element",POINTER(double_row_element),("col_index",c_int),("next_element),("col_index",c_int),("next_element),("col_index",c_int),("next_element),("col_index",c_int),("next_element),("next_element),("col_index",c_int),("next_element),("next_element),("next_element),("next_element),("next_element),("next_element),("next_element),("next_element),("next_element),("next_element),("next_element),("next_element),("next_element),("next_element),("next_element),("next_element),("next_element),("next_element),("next_element),("next_element),("next_element),("next_element),("next_element),("next_element),("next_element),("next_element),("next_element),("next_element),("next_element),("next_element),("next_element),("next_element),("next_element),("next_element),("next_element),("next_element),("next_element),("next_element),("next_element),("next_element),("next_element),("next_element),("next_element),("next_element),("next_element),("next_element),("next_element),("next_element),("next_element),("next_element),("next_element),("next_element),("next_element),("next_element),("next_element),("next_element),("next_element),("next_element),("next_element),("next_element),("next_element),("next_element),("next_element),("next_element),("next_element),("next_element),("next_element),("next_element),("next_element),("next_element),("next_element),("next_element),("next_element),("next_element),("next_element),("next_element),("next_element),("next_element),("next_element),("next_element),("next_element),("next_element),("next_element),("next_element),("next_element),("next_element),("next_element),("next_element),("next_element),("next_element),("next_element),("next_element),("next_element),("next_element),("next_element),("next_element),("next_element),("next_element),("
class double_sparse_matrix(Structure):
          _fields_=[("nrows",c_int),("ncols",c_int),("nnz",c_int),("rows",POINTER(POINTER(double_row_elements)
double_sparse_pointer=POINTER(double_sparse_matrix)
sparse_library=CDLL("/home/jkantor/linked_list_sparse.so")
\verb|initialize_matrix=sparse_library.initialize_matrix|\\
initialize_matrix.restype=double_sparse_pointer
set_value=sparse_library.set_value
get_value=sparse_library.get_value
get_value.restype=c_double
free_matrix=sparse_library.free_matrix
```

Lets discuss the above code. The original C code stored a sparse matrix as a linked list. The python code uses the ctypes Structure class to create structures mirroring the structs in the C code. To create python object representing a C struct, simply create class that derives from Structure. The _fields_ attribute of the class must be set to a list of tuples of field names and values. Note that in case you need to refer to a struct before it is completely defined (as in the linked list) you can first declare it with "Pass", and then specify the field contents as above. Also note the POINTER operator which creates a pointer out of any ctypes type. We are able to directly call our library as follows.

int i;

```
m=double_sparse_pointer()
m=initialize_matrix(c_int(10),c_int(10))
set_value(m,c_int(4),c_int(4),c_double(5.0))
a=get_value(m,c_int(4),c_int(4))
print("%f"%a)
free_matrix(m)
```

Note that you can access the contents of a structure just by (struct_object).field name. However for pointers, there is a contents attribute. So, in the above, m.contents.nrows would let you access the nrows field. In fact you can manually walk along the linked list as follows.

```
m=double_sparse_pointer()
m=initialize_matrix(c_int(10),c_int(10))
set_value(m,c_int(4),c_int(4),c_double(5.0))
a=m.contents.rows[4]
b=a.contents.next_element
b.contents.value
free matrix(m)
```

Comparison to Cython/Pyrex

is certainly possible to write a solver in Cython Pyrex. From the orhttp://www.scipy.org/PerformancePython?highlight=%28performance%29 website you can find an example. One potential downside to Cython over the previous solutions is it requires the user to understand how NumPy arrays or Sage matrices are implemented so as to be able to access their internal data. In contrast the weave, scipy, and ctypes examples only require the user to know C or Fortran and from their perspective the NumPy data magically gets passed to C or Fortran with no further thought from them. In order for pyrex to be competitive as a way to interactively write compiled code, the task of accessing the internal structure of NumPy arrays or Sage matrices needs to be hidden.

Parallel Computation

mpi4py

MPI which stands for message passing interface is a common library for parallel programming. There is a package mpi4py that builds on the top of mpi, and lets arbitrary python objects be passed between different processes. These packages are not part of the default sage install. To install them do

```
sage: optional_packages()
```

Find the package name openmpi-* and mpi4py-*and do

```
sage: install_package('openmpi-*')
sage: install_package('mpi4py-*')
```

Note that openmpi takes a while to compile (15-20 minutes or so). Openmpi can be run on a cluster, however this requires some set up so that processes on different machines can communicate (though if you are on a cluster this is probably already set up). The simplest case is if you are on a shared memory or multicore system where openmpi will just work with no configuration from you. To be honest, I have never tried to run mpi4py on a cluster, though there is much information about these topics online.

Now, the way that mpi works is you start a group of mpi processes, all of the processes run the same code. Each process has a rank, that is a number that identifies it. The following pseudocode indicates the general format of MPI programs.

```
if my rank is n:
   do some somputation ...
   send some stuff to the process of rank j
   receive some data from the process of rank k
else if my rank is n+1:
   ....
```

Each processes looks for what it's supposed to do (specified by its rank) and processes can send data and receive data. Lets give an example. Create a script with the following code in a file mpi_1.py

```
from mpi4py import MPI
comm = MPI.COMM_WORLD
print("hello world")
print("my rank is: %d"%comm.rank)
```

To run it you can do (from the command line in your sage directory)

```
./local/bin/mpirun -np 5 ./sage -python mpi_1.py
```

The command mpirun -np 5 starts 5 copies of a program under mpi. In this case we have 5 copies of sage in pure python mode run the script mpi_1.py. The result should be 5 "hello worlds" plus 5 distinct ranks. The two most important mpi operations are sending and receiving. Consider the following example which you should put in a script mpi_2.py

```
from mpi4py import MPI
import numpy
comm = MPI.COMM_WORLD
rank=comm.rank
size=comm.size
v=numpy.array([rank]*5,dtype=float)
comm.send(v,dest=(rank+1)%size)
data=comm.recv(source=(rank-1)%size)
print("my rank is %d"%rank)
print("I received this:")
print(data)
```

The same command as above with mpi_1.py replaced by mpi_2.py will produce 5 outputs and you will see each process creates an array and then passes it to the next guy (where the last guy passes to the first.) Note that MPI.size is the total number of mpi processes. MPI.COMM WORLD is the communication world.

There are some subtleties regarding MPI to be aware of. Small sends are buffered. This means if a process sends a small object it will be stored by openmpi and that process will continue its execution and the object it sent will be received whenever the destination executes a receive. However, if an object is large a process will hang until its destination executes a corresponding receive. In fact the above code will hang if [rank]*5 is replaced by [rank]*500. It would be better to do

```
from mpi4py import MPI
import numpy
comm = MPI.COMM_WORLD
rank=comm.rank
size=comm.size
v=numpy.array([rank]*500,dtype=float)
if comm.rank==0:
    comm.send(v,dest=(rank+1)%size)
if comm.rank > 0:
```

```
data=comm.recv(source=(rank-1)%size)
  comm.send(v,dest=(rank+1)%size)

if comm.rank==0:
    data=comm.recv(source=size-1)

print("my rank is %d"%rank)
print("I received this:")
print(data)
```

Now the first process initiates a send, and then process 1 will be ready to receive and then he will send and process 2 will be waiting to receive, etc. This will not lock regardless of how large of an array we pass.

A common idiom is to have one process, usually the one with rank 0 act as a leader. That processes sends data out to the other processes and processes the results and decides how further computation should proceed. Consider the following code

```
from mpi4py import MPI
import numpy
sendbuf=[]
root=0
comm = MPI.COMM_WORLD
if comm.rank==0:
    m=numpy.random.randn(comm.size,comm.size)
    print(m)
    sendbuf=m

v=comm.scatter(sendbuf,root)

print("I got this array:")
print(v)
```

The scatter command takes a list and evenly divides it amongst all the processes. Here the root process creates a matrix (which is viewed as a list of rows) and then scatters it to everybody (roots sendbuf is divided equally amongst the processes). Each process prints the row it got. Note that the scatter command is executed by everyone, but when root executes it, it acts as a send and a receive (root gets one row from itself), while for everyone else it is just a receive.

There is a complementary gather command that collects results from all the processes into a list. The next example uses scatter and gather together. Now the root process scatters the rows of a matrix, each process then squares the elements of the row it gets. Then the rows are all gathered up again by the root process who collects them into a new matrix.

```
from mpi4py import MPI
import numpy
comm = MPI.COMM_WORLD
sendbuf=[]
root=0
if comm.rank==0:
    m=numpy.array(range(comm.size*comm.size), dtype=float)
    m.shape=(comm.size,comm.size)
    print (m)
    sendbuf=m
v=comm.scatter(sendbuf,root)
print("I got this array:")
print(v)
\Delta\!=\!\Delta\!*\!\Delta
recvbuf=comm.gather(v,root)
if comm.rank==0:
```

```
print numpy.array(recvbuf)
```

There is also a broadcast command that sends a single object to every process. Consider the following small extension. This is the same as before, but now at the end the root process sends everyone the string "done", which is printed out.

```
v=MPI.COMM_WORLD.scatter(sendbuf,root)
print("I got this array:")
print(v)
v=v*v
recvbuf=MPI.COMM_WORLD.gather(v,root)
if MPI.COMM_WORLD.rank==0:
    print numpy.array(recvbuf)

if MPI.COMM_WORLD.rank==0:
    sendbuf="done"
recvbuf=MPI.COMM_WORLD.bcast(sendbuf,root)
print(recvbuf)
```

MPI programming is difficult. It is "schizophrenic programming" in that you are writing a single programming with multiple threads of execution "many voices in one head".

Parallel Laplace Solver

The following code solves Laplace's equation in parallel on a grid. It divides a square into n parallel strips where n is the number of processes and uses jacobi-iteration. The way the code works is that the root process creates a matrix and distributes the pieces to the other processes. At each iteration each process passes its upper row to the process above and its lower row to the process below since they need to know the values at neighboring points to do the iteration. Then they iterate and repeat. Every 500 iterations the error estimates from the processes are collected using Gather. you can compare the output of this with the solver we wrote in the section on f2py.

```
from mpi4py import MPI
import numpy
size=MPI.size
rank=MPI.rank
num_points=500
sendbuf=[]
root=0
dx=1.0/(num\_points-1)
from numpy import r_
j=numpy.complex(0,1)
rows_per_process=num_points/size
max_iter=5000
num_iter=0
total_err=1
def numpyTimeStep(u,dx,dy):
    dx2, dy2 = dx**2, dy**2
    dnr_inv = 0.5/(dx2 + dy2)
    u_old=u.copy()
    # The actual iteration
    u[1:-1, 1:-1] = ((u[0:-2, 1:-1] + u[2:, 1:-1])*dy2 +
                      (u[1:-1,0:-2] + u[1:-1, 2:])*dx2)*dnr_inv
    v = (u - u_old).flat
    return u, numpy.sqrt(numpy.dot(v, v))
```

```
if MPI.rank==0:
   print("num_points: %d"%num_points)
   print("dx: %f"%dx)
    print("row_per_procss: %d"%rows_per_process)
    m=numpy.zeros((num_points,num_points),dtype=float)
   pi_c=numpy.pi
    x=r_[0.0:pi_c:num_points*j]
   m[0,:]=numpy.sin(x)
   m[num_points-1,:]=numpy.sin(x)
    l=[ m[i*rows_per_process:(i+1)*rows_per_process,:] for i in range(size)]
    sendbuf=1
my_grid=MPI.COMM_WORLD.Scatter(sendbuf,root)
while num_iter < max_iter and total_err > 10e-6:
    if rank==0:
        MPI.COMM_WORLD.Send(my_grid[-1,:],1)
    if rank > 0 and rank< size-1:</pre>
        row_above=MPI.COMM_WORLD.Recv(rank-1)
        MPI.COMM_WORLD.Send(my_grid[-1,:],rank+1)
    if rank==size-1:
        row_above=MPI.COMM_WORLD.Recv(MPI.rank-1)
        MPI.COMM_WORLD.Send(my_grid[0,:],rank-1)
    if rank > 0 and rank< size-1:</pre>
        row_below=MPI.COMM_WORLD.Recv(MPI.rank+1)
        MPI.COMM_WORLD.Send(my_grid[0,:],MPI.rank-1)
    if rank==0:
        row_below=MPI.COMM_WORLD.Recv(1)
    if rank >0 and rank < size-1:</pre>
        row_below.shape=(1,num_points)
        row_above.shape=(1,num_points)
        u,err =numpyTimeStep(r_[row_above,my_grid,row_below],dx,dx)
        my_grid=u[1:-1,:]
    if rank==0:
        row_below.shape=(1,num_points)
        u,err=numpyTimeStep(r_[my_grid,row_below],dx,dx)
        my_grid=u[0:-1,:]
    if rank==size-1:
        row_above.shape=(1, num_points)
        u,err=numpyTimeStep(r_[row_above,my_grid],dx,dx)
        my_grid=u[1:,:]
    if num_iter%500==0:
        err_list=MPI.COMM_WORLD.Gather(err,root)
        if rank==0:
```

For small grid sizes this will be slower than a straightforward serial implementation, this is because there is overhead from the communication, and for small grids the interprocess communication takes more time than just doing the iteration. However, on a 1000x1000 grid I find that using 4 processors, the parallel version takes only 6 seconds while the serial version we wrote earlier takes 20 seconds.

Excercise: Rewrite the above using f2py or weave, so that each process compiles a fortran or C timestep function and uses that, how fast can you get this?

Visualization

One of the most common uses of computer algebra systems is of course plotting. At the moment Sage does not have much in the way of intrinsic support for visualization. However, there are some very powerful open source programs for visualization that you can use from within Sage. The goal of this chapter is to explain how to set them up. The main tool we will be using is VTK, http://www.vtk.org. VTK is an amazing visualization tool, and the results that can be achieved are incredible. It is probably one of the best visualization tools in existence, and can probably do anything you might want a visualization tool to do. However, because it can do so much, it is a bit tricky to use potentially. Luckily there are a few tools for using VTK from within python that make it easier to get started. The ones we will focus on are MavaVi, http://mayavi.sourceforge.net/, and easyviz.

Installation of Visualization Tools

Installing Visualization Tools on Linux This section assumes you are running linux. You may need administrator rights to complete this section. First try

```
sage: import Tkinter
```

If this works great if not it means your sage was not compiled with tcl/tk bindings. There are two possible reasons. If you used a binary then this will be the case. If you built from source but don't have the tcl/tk libraries on your system you will have the same problem. To fix this install the tcl and tk libraries and development (source and headers) packages for your linux distribution. Now you need to rebuild Sage's python

```
sage: install_package('python-2.5<version>.spkg')
```

where you should replace < version > by the version of python. Type

```
sage: !ls spkg/standard | grep python-2.5
```

This will give you the name of the python package to use above. In the case that it gives multiple names, choose the one with the highest version number. Now again try

```
sage: import Tkinter
```

If this works we can install vtk, but first you need cmake. Test if your system has it by typing cmake at the shell, (or !cmake in sage). If cmake is on your system then you are ready. If not either install cmake on your system using your distributions tools or do

```
sage: install_package('cmake-2.4.7')
```

Now we want to compile VTK which does all the hard work. To do this make sure you have the opengl libraries for you system installed. This will be something like libgl1-mesa-glx, libgl1-mesa-dev.

```
sage: install_package('vtk-5.0.3.p1')
```

This will take quite a while to compile, probably 20 min to an hour or so. Once this is done we install the python wrappers, the next part takes about 10 seconds,

```
sage: install_package('MayaVi-1.5')
sage: install_package('scitools++')
sage: install_package('PyVTK-0.4.74')
```

Now you're done.

Installing Visualization Software on OS X The first thing we need to do is rebuild Python to use OSX's frameworks, so that it can create graphical windows. To do this first from the terminal do

```
cd $SAGE_ROOT/local/lib
rm libpng*.dylib
```

where \$SAGE_ROOT is the directory of your Sage install. Next from within Sage,

```
sage: install_package('python-2.5.1-framework')
```

Next we will build vtk, this will take a while

```
sage: install_package('vtk-5.0.3.p1')

Finally

sage: install_package('MayaVi-1.5')
sage: install_package('scitools++')
sage: install_package('PyVTK-0.4.74')
```

Plotting

We will plot a surface two ways. First we will use easyviz. Consider the following code:

```
import numpy
from scitools import easyviz
x = numpy.arange(-8,8,.2)
```

```
xx,yy = numpy.meshgrid(x,x)
r = numpy.sqrt(xx**2+yy**2) + 0.01
zz = numpy.sin(r)/r
easyviz.surfc(x,x,zz)
```

The function surfc takes a list of x coordinates, and y coordinates and a numpy array z. Its plots a surface that has height z[i,j] at the point (x[i],y[i]). Note the use of meshgrid, and vectorized numpy functions that let us evaluate $\frac{\sin(\sqrt{x^2+y^2})+1}{\sqrt{x^2+y^2}+1}$ over the grid very easily. We discussed meshgrid at the beginning when we were talking about numpy. Note that you can drag the plot around with your mouse and look at it from different angles.

We can make this plot look a bit nicer by adding some shading and nicer coloring and some labels as follows.

Let us now try to plot some vector fields. Consider the following code

```
import numpy
from scitools import easyviz
RealNumber=float
Integer=int
j=numpy.complex(0,1)
w=numpy.zeros((5,5,5))
u=w+1.0
xx,yy,zz=numpy.mgrid[-1.0:1.0:5*j,-1:1:5*j,-1:1:5*j]
easyviz.quiver3(xx,yy,zz,w,w,u)
```

This should plot a vector field that points up everywhere. The arguments to quiver are 6, $n \times n \times n$ arrays. The first three arrays are the location of the vectors, that is there will be a vector at (xx[i,j,k],yy[i,j,k],zz[i,j,k]) for $0 \le i,j,k < n$. The second three arrays are the directions, i.e., the vector at (xx[i,j,k],yy[i,j,k],zz[i,j,k]) points in the direction (w[i,j,k],w[i,j,k],u[i,j,k]).

Now let us give some examples with MayaVi. First lets see how to plot a function like we did with easyviz.

```
import numpy
from mayavi.tools import imv
x=numpy.arange(-8,8,.2)
def f(x,y):
    r=numpy.sqrt(x**2+y**2)+.01
    return numpy.sin(r)/r
imv.surf(x,x,f)
```

This will open mayavi, and display the plot of the function. The first two arguments to surf are arrays x and y, s.t. the function will be evaluated at (x[i], y[j]). The last argument is the function to graph. It probably looks a bit different

than the easyviz example. Lets try to make it look similar to the easyviz example. First note that on the left there is a list of filters and modules. Double-click the warpscalars button in the filters menu, and change the scale factor from 1 to say 5. This should redraw the graph similar to how easyviz drew it. There are quite a few other options you can play around with. For example, next click on the module surfacemap, and you will see you can make the graph transparent by changing the opacity. You can also change it to a wireframe or make it plot contours.

TODO: More examples

12.1.18 Three Lectures about Explicit Methods in Number Theory Using Sage

This article is about using the mathematical software Sage to do computations with number fields and modular forms. It was written for the October 2008 Bordeaux meeting on explicit methods in number theory (http://www.math.u-bordeaux.fr/gtem2008/). It assumes no prior knowledge about Sage, but assumes a graduate level background in algebraic number theory.

Introduction

What is Sage?

Sage (see http://sagemath.org) is a comprehensive mathematical software system for computations in many areas of pure and applied mathematics. We program Sage using the mainstream programming language Python (see http://python.org), or its compiled variant Cython. It is also very easy to efficiently use code written in C/C++ from Sage.

The author of this article started the Sage project in 2005.

Sage is free and open source, meaning you can change any part of Sage and redistribute the result without having to pay any license fees, and Sage can also leverage the power of commercial mathematical software such as Magma and Mathematica, if you happen to have access to those closed source commercial systems.

This document assumes no prior knowledge of either Python or Sage. Our goal is to help number theorists do computations involving number fields and modular forms using Sage.

TODO: Overview of Article

As you read this article, please try every example in Sage, and make sure things works as I claim, and do all of the exercises. Moreover, you should experiment by typing in similar examples and checking that the output you get agrees with what you expect.

Using Sage

To use Sage, install it on your computer, and use either the command line or start the Sage notebook by typing notebook () at the command line.

We show Sage sessions as follows:

```
sage: factor(123456)
2^6 * 3 * 643
```

This means that if you type factor (123456) as input to Sage, then you'll get $2^6 * 3 * 643$ as output. If you're using the Sage command line, you type factor (123456) and press enter; if you're using the Sage notebook via your web browser, you type factor (123456) into an input cell and press shift-enter; in the output cell you'll see $2^6 * 3 * 643$.

After trying the factor command in the previous paragraph (do this now!), you should try factoring some other numbers.

Note: What happens if you factor a negative number? a rational number?

You can also draw both 2d and 3d pictures using Sage. For example, the following input plots the number of prime divisors of each positive integer up to 500.

```
sage: line([(n, len(factor(n))) for n in [1..500]])
Graphics object consisting of 1 graphics primitive
```

And, this example draws a similar 3d plot:

```
sage: import warnings
sage: warnings.simplefilter('ignore', UserWarning)
sage: v = [[len(factor(n*m)) for n in [1..15]] for m in [1..15]]
sage: list_plot3d(v, interpolation_type='nn')
Graphics3d Object
```

The Sage-Pari-Magma Ecosystem

- The main difference between Sage and Pari is that Sage is vastly larger than Pari with a much wider range of functionality, and has many more data types and much more structured objects. Sage in fact includes Pari, and a typical Sage install takes nearly a gigabyte of disk space, whereas a typical Pari install is much more nimble, using only a few megabytes. There are many number-theoretic algorithms that are included in Sage, which have never been implemented in Pari, and Sage has 2d and 3d graphics which can be helpful for visualizing number theoretic ideas, and a graphical user interface. Both Pari and Sage are free and open source, which means anybody can read or change anything in either program, and the software is free.
- The biggest difference between Sage and Magma is that Magma is closed source, not free, and difficult for users to extend. This means that most of Magma cannot be changed except by the core Magma developers, since Magma itself is well over two million lines of compiled C code, combined with about a half million lines of interpreted Magma code (that anybody can read and modify). In designing Sage, we carried over some of the excellent design ideas from Magma, such as the parent, element, category hierarchy.
- Any mathematician who is serious about doing extensive computational work in algebraic number theory and
 arithmetic geometry is strongly urged to become familiar with all three systems, since they all have their pros and
 cons. Pari is sleek and small, Magma has much unique functionality for computations in arithmetic geometry,
 and Sage has a wide range of functionality in most areas of mathematics, a large developer community, and
 much unique new code.

Number Fields

Introduction to Number Fields

In Sage, we can create the number field $\mathbf{Q}(\sqrt[3]{2})$ as follows.

```
sage: K.<alpha> = NumberField(x^3 - 2)
```

The above creates *two* Sage objects, K and α . Here K "is" (isomorphic to) the number field $\mathbf{Q}(\sqrt[3]{2})$, as we confirm below:

and α is a root of x^3-2 , so α is an abstract choice of $\sqrt[3]{2}$ (no specific embedding of the number field K into C is chosen by default in Sage-3.1.2):

```
sage: alpha^3
2
sage: (alpha+1)^3
3*alpha^2 + 3*alpha + 3
```

The variable x Note that we did *not* define x above before using it. You could "break" the above example by redefining x to be something funny:

```
sage: x = 1
sage: K.<alpha> = NumberField(x^3 - 2)
Traceback (most recent call last):
...
TypeError: polynomial (=-1) must be a polynomial.
```

The *Traceback* above indicates that there was an error. Potentially lots of detailed information about the error (a "traceback") may be given after the word Traceback and before the last line, which contains the actual error messages.

Note: *Important*: whenever you use Sage and get a big error, look at the last line for the actual error, and only look at the rest if you are feeling adventurous. In the notebook, the part indicated by . . . above is not displayed; to see it, click just to the left of the word *Traceback* and the traceback will appear.

If you redefine x as above, but need to define a number field using the indeterminate x, you have several options. You can reset x to its default value at the start of Sage, you can redefine x to be a symbolic variable, or you can define x to be a polynomial indeterminate (a polygen):

```
sage: reset('x')
sage: x
x
sage: x = 1
sage: x = var('x')
sage: x
x
sage: x = 1
sage: x = polygen(QQ, 'x')
sage: x
x
sage: x = 1
sage: R.<x> = PolynomialRing(QQ)
sage: x
x
```

Using tab completion to get the methods of an object One you have created a number field K, type K. [tab key] to see a list of functions. Type, e.g., K.Minkowski_embedding?[tab key] to see help on the Minkowski_embedding command. To see source code, type K.Minkowski_embedding??[tab key].

```
sage: K.<alpha> = NumberField(x^3 - 2)
sage: K.[tab key]
```

Symbolic Expressions Another natural way for us to create certain number fields is to create a symbolic expression and adjoin it to the rational numbers. Unlike Pari and Magma (and like Mathematica and Maple), Sage also supports

manipulation of symbolic expressions and solving equations, without defining abstract structures such as a number fields. For example, we can define a variable $a=\sqrt{2}$ as an abstract symbolic object by simply typing $a=\operatorname{sqrt}(2)$. When we type parent (a) below, Sage tells us the mathematical object that it views a as being an element of; in this case, it's the ring of all symbolic expressions.

```
sage: a = sqrt(2)
sage: parent(a)
Symbolic Ring
```

sqrt(2) in Pari and Magma In particular, typing sqrt(2) does *not* numerically extract an approximation to $\sqrt{2}$, like it would in Pari or Magma. We illustrate this below by calling Pari (via the gp interpreter) and Magma directly from within Sage. After we evaluate the following two input lines, copies of GP/Pari and Magma are running, and there is a persistent connection between Sage and those sessions.

```
sage: gp('sqrt(2)')
1.414213562373095048801688724...
sage: magma('Sqrt(2)') # optional - magma
1.414213562373095048801688724...
```

You probably noticed a pause when evaluated the second line as Magma started up. Also, note the # optional comment, which indicates that the line won't work if you don't have Magma installed.

Numerically evaluating sqrt(2) Incidentally, if you want to numerically evaluate $\sqrt{2}$ in Sage, just give the optional prec argument to the sqrt function, which takes the required number of *bits* (binary digits) of precision.

```
sage: sqrt(2, prec=100)
1.4142135623730950488016887242
```

It's important to note in computations like this that there is not an *a priori* guarantee that prec bits of the *answer* are all correct. Instead, what happens is that Sage creates the number 2 as a floating point number with 100 bits of accuracy, then asks Paul Zimmerman's MPFR C library to compute the square root of that approximate number.

Arithmetic with sqrt(2) We return now to our symbolic expression $a = \sqrt{2}$. If you ask to square a + 1 you simply get the formal square. To expand out this formal square, we use the expand command.

```
sage: a = sqrt(2)
sage: (a+1)^2
(sqrt(2) + 1)^2
sage: expand((a+1)^2)
2*sqrt(2) + 3
```

Adjoining a symbolic expression Given any symbolic expression for which Sage can compute its minimal polynomial, you can construct the number field obtained by adjoining that expression to \mathbf{Q} . The notation is quite simple - just type QQ[a] where a is the symbolic expression.

```
sage: a = sqrt(2)
sage: K. <b> = QQ[a]
sage: K
Number Field in sqrt2 with defining polynomial x^2 - 2
sage: b
sqrt2
sage: (b+1)^2
2*sqrt2 + 3
```

```
sage: QQ[a/3 + 5] Number Field in a with defining polynomial x^2 - 10*x + 223/9
```

Coercion: QQ[a] versus QQ(a) You can't create the number field **Q**(a) in Sage by typing QQ (a), which has a *very different* meaning in Sage. It means "try to create a rational number from a." Thus QQ (a) in Sage is the analogue of QQ! a in Magma (Pari has no notion of rings such as QQ).

```
sage: a = sqrt(2)
sage: QQ(a)
Traceback (most recent call last):
...
TypeError: unable to convert sqrt(2) to a rational
```

In general, if X is a ring, or vector space or other "parent structure" in Sage, and a is an element, type X (a) to make an element of X from a. For example, if X is the finite field of order 7, and a = 2/5 is a rational number, then X (a) is the finite field element 6 (as a quick exercise, check that this is mathematically the correct interpretation).

```
sage: X = GF(7); a = 2/5
sage: X(a)
```

Solving a cubic equation As a slightly less trivial illustration of symbolic manipulation, consider the cubic equation

$$x^3 + \sqrt{2}x + 5 = 0.$$

In Sage, we can create this equation, and find an exact symbolic solution.

```
sage: x = var('x')

sage: eqn = x^3 + sqrt(2) *x + 5 == 0

sage: a = solve(eqn, x)[0].rhs()
```

The first line above makes sure that the symbolic variable x is defined, the second creates the equation eqn, and the third line solves eqn for x, extracts the first solution (there are three), and takes the right hand side of that solution and assigns it to the variable a.

Viewing complicated symbolic expressions To see the solution nicely typeset, use the pretty_print command

```
sage: pretty_print(a)
<html><script type="math/tex">\newcommand{\Bold}[1]{\mathbf{#1}}-\frac{1}{2} ...
```

$$-\frac{1}{2} \left(i \sqrt{3}+1\right) \left(\frac{1}{18} \sqrt{8 \sqrt{2}+675} \sqrt{3}-\frac{5}{2}\right)^{\left(\frac{1}{3}\right)}+\frac{1}{6} \frac{(-i \sqrt{3}+1) \sqrt{2}}{\left(\frac{1}{18} \sqrt{8 \sqrt{2}+675} \sqrt{3}-\frac{5}{2}\right)^{\left(\frac{1}{3}\right)}}$$

You can also see the latex needed to paste a into a paper by typing latex (a). The latex command works on most Sage objects.

```
sage: latex(a)
-\frac{1}{2} \, {\left(i \, \sqrt{3} + 1\right)} ...
```

Adjoining a root of the cubic Next, we construct the number field obtained by adjoining the solution a to **Q**. Notice that the minimal polynomial of the root is $x^6 + 10x^3 - 2x^2 + 25$.

Warning: The following tests are currently broken until Trac ticket #5338 is fixed.

```
sage: K.<b> = QQ[a]
sage: K
Number Field in a with defining
polynomial x^6 + 10*x^3 - 2*x^2 + 25
sage: a.minpoly()
x^6 + 10*x^3 - 2*x^2 + 25
sage: b.minpoly()
x^6 + 10*x^3 - 2*x^2 + 25
```

We can now compute interesting invariants of the number field ${\cal K}$

```
sage: K.class_number()
5
sage: K.galois_group().order()
72
```

Number Fields: Galois Groups and Class Groups

Galois Groups We can compute the Galois group of a number field using the galois_group function, which by default calls Pari (http://pari.math.u-bordeaux.fr/). You do not have to worry about installing Pari, since *Pari is part of Sage*. In fact, despite appearances much of the difficult algebraic number theory in Sage is actually done by the Pari C library (be sure to also cite Pari in papers that use Sage).

```
sage: K.<alpha> = NumberField(x^6 + 40*x^3 + 1372)
sage: G = K.galois_group()
sage: G
Galois group of Number Field in alpha with defining polynomial x^6 + 40*x^3 + 1372
```

Internally G is represented as a group of permutations, but we can also apply any element of G to any element of the field:

```
sage: G.order()
6
sage: G.gens()
[(1,2)(3,4)(5,6), (1,4,6)(2,5,3)]
sage: f = G.1; f(alpha)
1/36*alpha^4 + 1/18*alpha
```

Some more advanced number-theoretical tools are available via G:

```
sage: P = K.primes_above(2)[0]
sage: G.inertia_group(P)
Subgroup [(), (1,4,6)(2,5,3), (1,6,4)(2,3,5)] of Galois group of Number Field in alpha with defining
sage: sorted([G.artin_symbol(Q) for Q in K.primes_above(5)]) # random order, see Trac #18308
[(1,3)(2,6)(4,5), (1,2)(3,4)(5,6), (1,5)(2,4)(3,6)]
```

If the number field is not Galois over \mathbf{Q} , then the <code>galois_group</code> command will construct its Galois closure and return the Galois group of that; you need to give it a variable name for the generator of the Galois closure:

```
sage: K.<a> = NumberField(x^3 - 2)
sage: G = K.galois_group(names='b'); G
Galois group of Galois closure in b of Number Field in a with defining polynomial x^3 - 2
sage: G.order()
```

Some more Galois groups We compute two more Galois groups of degree 5 extensions, and see that one has Galois group S_5 , so is not solvable by radicals. For these purposes we only want to know the structure of the Galois group as an abstract group, rather than as an explicit group of automorphisms of the splitting field; this is much quicker to calculate. PARI has a type for representing "abstract Galois groups", and Sage can use this.:

```
sage: NumberField(x^5 - 2, 'a').galois_group(type="pari")
Galois group PARI group [20, -1, 3, "F(5) = 5:4"] of
degree 5 of the Number Field in a with defining
polynomial x^5 - 2
sage: NumberField(x^5 - x + 2, 'a').galois_group(type="pari")
Galois group PARI group [120, -1, 5, "S5"] of degree 5 of
the Number Field in a with defining polynomial x^5 - x + 2
```

Magma's Galois group command Recent versions of Magma have an algorithm for computing Galois groups that in theory applies when the input polynomial has any degree. There are no open source implementation of this algorithm (as far as I know). If you have Magma, you can use this algorithm from Sage by calling the <code>galois_group</code> function and giving the <code>algorithm='magma'</code> option. The return value is one of the groups in the GAP transitive groups database.

```
sage: K.<a> = NumberField(x^3 - 2)
sage: K.galois_group(type="gap", algorithm='magma') # optional - magma database_gap
Galois group Transitive group number 2 of degree 3 of
the Number Field in a with defining polynomial x^3 - 2
```

We emphasize that the above example should not work if you don't have Magma.

Computing complex embeddings You can also enumerate all complex embeddings of a number field:

```
sage: K.complex_embeddings()
[
Ring morphism:
    From: Number Field in a with defining polynomial x^3 - 2
    To:    Complex Field with 53 bits of precision
    Defn: a |--> -0.629960524947437 - 1.09112363597172*I,
Ring morphism:
    From: Number Field in a with defining polynomial x^3 - 2
    To:    Complex Field with 53 bits of precision
    Defn: a |--> -0.629960524947437 + 1.09112363597172*I,
Ring morphism:
    From: Number Field in a with defining polynomial x^3 - 2
    To:    Complex Field with 53 bits of precision
    Defn: a |--> 1.25992104989487
]
```

Class Numbers and Class Groups The class group C_K of a number field K is the group of fractional ideals of the maximal order R of K modulo the subgroup of principal fractional ideals. One of the main theorems of algebraic

number theory asserts that C_K is a finite group. For example, the quadratic number field $\mathbf{Q}(\sqrt{-23})$ has class number 3, as we see using the Sage class number command.

```
sage: L.<a> = NumberField(x^2 + 23)
sage: L.class_number()
3
```

Quadratic imaginary fields with class number 1 There are only 9 quadratic imaginary field $\mathbf{Q}(\sqrt{D})$ that have class number 1:

$$D = -3, -4, -7, -8, -11, -19, -43, -67, -163$$

To find this list using Sage, we first experiment with making lists in Sage. For example, typing [1..10] makes the list of integers between 1 and 10.

```
sage: [1..10]
[1, 2, 3, 4, 5, 6, 7, 8, 9, 10]
```

We can also make the list of odd integers between 1 and 11, by typing [1, 3, ..., 11], i.e., by giving the second term in the arithmetic progression.

```
sage: [1,3,..,11]
[1, 3, 5, 7, 9, 11]
```

Applying this idea, we make the list of negative numbers from -1 down to -10.

```
sage: [-1,-2,..,-10] [-1, -2, -3, -4, -5, -6, -7, -8, -9, -10]
```

Enumerating quadratic imaginary fields with class number 1 The first two lines below makes a list v of every D from -1 down to -200 such that D is a fundamental discriminant (the discriminant of a quadratic imaginary field).

Note: Note that you will not see the ... in the output below; this ... notation just means that part of the output is omitted below.

```
sage: w = [-1, -2, ..., -200]

sage: v = [D \text{ for } D \text{ in } w \text{ if } is\_fundamental\_discriminant(D)]

sage: v

[-3, -4, -7, -8, -11, -15, -19, -20, ..., -195, -199]
```

Finally, we make the list of D in our list v such that the quadratic number field $\mathbf{Q}(\sqrt{D})$ has class number 1. Notice that QuadraticField(D) is a shorthand for NumberField(x^2 - D).

```
sage: [D for D in v if QuadraticField(D,'a').class_number()==1]
[-3, -4, -7, -8, -11, -19, -43, -67, -163]
```

Of course, we have *not* proved that this is the list of all negative D so that $\mathbf{Q}(\sqrt{D})$ has class number 1.

Class number 1 fields A frustrating open problem is to prove that there are infinitely many number fields with class number 1. It is quite easy to be convinced that this is probably true by computing a bunch of class numbers of real quadratic fields. For example, over 58 percent of the real quadratic number fields with discriminant D < 1000 have class number 1!

```
sage: w = [1..1000]
sage: v = [D for D in w if is_fundamental_discriminant(D)]
sage: len(v)
302
sage: len([D for D in v if QuadraticField(D,'a').class_number() == 1])
176
sage: 176.0/302
0.582781456953642
```

For more intuition about what is going on, read about the Cohen-Lenstra heuristics.

Class numbers of cyclotomic fields Sage can also compute class numbers of extensions of higher degree, within reason. Here we use the shorthand CyclotomicField (n) to create the number field $\mathbf{Q}(\zeta_n)$.

```
sage: CyclotomicField(7)
Cyclotomic Field of order 7 and degree 6
sage: for n in [2..15]: print n, CyclotomicField(n).class_number()
2 1
3 1
...
15 1
```

In the code above, the notation for n in [2..15]: ... means "do ... for n equal to each of the integers $2, 3, 4, \ldots, 15$."

Note: Exercise: Compute what is omitted (replaced by ...) in the output of the previous example.

Assuming conjectures to speed computations Computations of class numbers and class groups in Sage is done by the Pari C library, and *unlike in Pari*, by default Sage tells Pari *not to assume* any conjectures. This can make some commands vastly slower than they might be directly in Pari, which *does assume unproved conjectures* by default. Fortunately, it is easy to tell Sage to be more permissive and allow Pari to assume conjectures, either just for this one call or henceforth for all number field functions. For example, with proof=False it takes only a few seconds to verify, modulo the conjectures assumed by Pari, that the class number of $\mathbf{Q}(\zeta_{23})$ is 3.

```
sage: CyclotomicField(23).class_number(proof=False)
3
```

Note: Exercise: What is the smallest n such that $\mathbf{Q}(\zeta_n)$ has class number bigger than 1?

Class group structure In addition to computing class numbers, Sage can also compute the group structure and generators for class groups. For example, the quadratic field $\mathbf{Q}(\sqrt{-30})$ has class group $C = (\mathbf{Z}/2\mathbf{Z})^{\oplus 2}$, with generators the ideal classes containing $(5, \sqrt{-30})$ and $(3, \sqrt{-30})$.

```
sage: K.<a> = QuadraticField(-30)
sage: C = K.class_group()
sage: C
Class group of order 4 with structure C2 x C2 of Number Field
in a with defining polynomial x^2 + 30
sage: category(C)
Category of finite commutative groups
sage: C.gens()
(Fractional ideal class (2, a), Fractional ideal class (3, a))
```

```
sage: K.<a> = QuadraticField(-30)
sage: C = K.class_group()
sage: C.0
Fractional ideal class (2, a)
sage: C.0.ideal()
Fractional ideal (2, a)
sage: I = C.0 * C.1
sage: I
Fractional ideal class (5, a)
```

Next we find that the class of the fractional ideal $(2, \sqrt{-30} + 4/3)$ is equal to the ideal class C.0.

```
sage: A = K.ideal([2, a+4/3])
sage: J = C(A)
sage: J
Fractional ideal class (2/3, 1/3*a)
sage: J == C.0
True
```

Unfortunately, there is currently no Sage function that writes a fractional ideal class in terms of the generators for the class group.

Orders and Relative Extensions

Orders in Number Fields An *order* in a number field K is a subring of K whose rank over \mathbb{Z} equals the degree of K. For example, if $K = \mathbb{Q}(\sqrt{-1})$, then $\mathbb{Z}[7i]$ is an order in K. A good first exercise is to prove that every element of an order is an algebraic integer.

```
sage: K.<I> = NumberField(x^2 + 1)
sage: R = K.order(7*I)
sage: R
Order in Number Field in I with defining polynomial x^2 + 1
sage: R.basis()
[1, 7*I]
```

Using the discriminant command, we compute the discriminant of this order

```
sage: factor(R.discriminant())
-1 * 2^2 * 7^2
```

Constructing the order with given generators You can give any list of elements of the number field, and it will generate the smallest ring R that contains them.

```
sage: K.<a> = NumberField(x^4 + 2)
sage: K.order([12*a^2, 4*a + 12]).basis()
[1, 4*a, 4*a^2, 16*a^3]
```

If R isn't of rank equal to the degree of the number field (i.e., R isn't an order), then you'll get an error message.

```
sage: K.order([a^2])
Traceback (most recent call last):
```

```
...
ValueError: the rank of the span of gens is wrong
```

Computing Maximal Orders We can also compute the maximal order, using the maximal order command, which behind the scenes finds an integral basis using Pari's nfbasis command. For example, $\mathbf{Q}(\sqrt[4]{2})$ has maximal order $\mathbf{Z}[\sqrt[4]{2}]$, and if α is a root of $x^3 + x^2 - 2x + 8$, then $\mathbf{Q}(\alpha)$ has maximal order with **Z**-basis

$$1, \frac{1}{2}a^2 + \frac{1}{2}a, a^2.$$

```
sage: K.<a> = NumberField(x^4 + 2)
sage: K.maximal_order().basis()
[1, a, a^2, a^3]
sage: L.<a> = NumberField(x^3 + x^2 - 2*x+8)
sage: L.maximal_order().basis()
[1, 1/2*a^2 + 1/2*a, a^2]
sage: L.maximal_order().basis()[1].minpoly()
x^3 - 2*x^2 + 3*x - 10
```

Functionality for non-maximal orders is minimal There is still much important functionality for computing with non-maximal orders that is missing in Sage. For example, there is no support at all in Sage for computing with modules over orders or with ideals in non-maximal orders.

```
sage: K.<a> = NumberField(x^3 + 2)
sage: R = K.order(3*a)
sage: R.ideal(5)
Traceback (most recent call last):
...
NotImplementedError: ideals of non-maximal orders not
yet supported.
```

Relative Extensions A *relative number field* L is a number field of the form $K(\alpha)$, where K is a number field, and an *absolute number field* is a number field presented in the form $\mathbf{Q}(\alpha)$. By the primitive element theorem, any relative number field $K(\alpha)$ can be written as $\mathbf{Q}(\beta)$ for some $\beta \in L$. However, in practice it is often convenient to view L as $K(\alpha)$. In *Symbolic Expressions*, we constructed the number field $\mathbf{Q}(\sqrt{2})(\alpha)$, where α is a root of $x^3 + \sqrt{2}x + 5$, but *not* as a relative field—we obtained just the number field defined by a root of $x^6 + 10x^3 - 2x^2 + 25$.

Constructing a relative number field step by step To construct this number field as a relative number field, first we let K be $\mathbf{Q}(\sqrt{2})$.

```
sage: K.<sqrt2> = QuadraticField(2)
```

Next we create the univariate polynomial ring R = K[X]. In Sage, we do this by typing R < X > = K[]. Here R < X > means "create the object R with generator X" and K[] means a "polynomial ring over K", where the generator is named based on the aformentioned X (to create a polynomial ring in two variables X, Y simply replace R < X > by R < X, Y >).

```
sage: R.<X> = K[]
sage: R
Univariate Polynomial Ring in X over Number Field in sqrt2
with defining polynomial x^2 - 2
```

Now we can make a polynomial over the number field $K = \mathbf{Q}(\sqrt{2})$, and construct the extension of K obtained by adjoining a root of that polynomial to K.

```
sage: L.<a> = K.extension(X^3 + sqrt2*X + 5)
sage: L
Number Field in a with defining polynomial X^3 + sqrt2*X + 5...
```

Finally, L is the number field $\mathbf{Q}(\sqrt{2})(\alpha)$, where α is a root of $X^3 + \sqrt{2}\alpha + 5$. We can do now do arithmetic in this number field, and of course include $\sqrt{2}$ in expressions.

```
sage: a^3
-sqrt2*a - 5
sage: a^3 + sqrt2*a
-5
```

Functions on relative number fields The relative number field L also has numerous functions, many of which have both relative and absolute version. For example the relative_degree function on L returns the relative degree of L over K; the degree of L over R is given by the absolute_degree function. To avoid possible ambiguity degree is not implemented for relative number fields.

```
sage: L.relative_degree()
3
sage: L.absolute_degree()
6
```

Extra structure on relative number fields Given any relative number field you can also an absolute number field that is isomorphic to it. Below we create $M = \mathbf{Q}(b)$, which is isomorphic to L, but is an absolute field over \mathbf{Q} .

```
sage: M.<b> = L.absolute_field()
sage: M
Number Field in b with defining
polynomial x^6 + 10*x^3 - 2*x^2 + 25
```

The structure function returns isomorphisms in both directions between M and L.

```
sage: M.structure()
(Isomorphism map:
    From: Number Field in b with defining polynomial x^6 + 10*x^3 - 2*x^2 + 25
    To: Number Field in a with defining polynomial X^3 + sqrt2*X + 5 over its base field, Isomorphism
    From: Number Field in a with defining polynomial X^3 + sqrt2*X + 5 over its base field
    To: Number Field in b with defining polynomial x^6 + 10*x^3 - 2*x^2 + 25)
```

Arbitrary towers of relative number fields In Sage one can create arbitrary towers of relative number fields (unlike in Pari, where a relative extension must be a single extension of an absolute field).

```
sage: R.<X> = L[]
sage: Z.<b> = L.extension(X^3 - a)
sage: Z
Number Field in b with defining polynomial X^3 - a over its base field
sage: Z.absolute_degree()
18
```

Note: Exercise: Construct the relative number field $L = K(\sqrt[3]{\sqrt{2} + \sqrt{3}})$, where $K = \mathbb{Q}(\sqrt{2}, \sqrt{3})$.

Relative number field arithmetic can be slow One shortcoming with relative extensions in Sage is that behind the scenes all arithmetic is done in terms of a single absolute defining polynomial, and in some cases this can be very slow (much slower than Magma). Perhaps this could be fixed by using Singular's multivariate polynomials modulo an appropriate ideal, since Singular polynomial arithmetic is extremely fast. Also, Sage has very little direct support for constructive class field theory, which is a major motivation for explicit computation with relative orders; it would be good to expose more of Pari's functionality in this regard.

A Bird's Eye View

We now take a whirlwind tour of some of the number theoretical functionality of Sage. There is much that we won't cover here, but this should help give you a flavor for some of the number theoretic capabilities of Sage, much of which is unique to Sage.

Integer Factorization

Quadratic Sieve Bill Hart's quadratic sieve is included with Sage. The quadratic sieve is the best algorithm for factoring numbers of the form pq up to around 100 digits. It involves searching for relations, solving a linear algebra problem modulo 2, then factoring n using a relation $x^2 \equiv y^2 \mod n$.

```
sage: qsieve(next_prime(2^90)*next_prime(2^91), time=True) # not tested
([1237940039285380274899124357, 2475880078570760549798248507],
'14.94user 0.53system 0:15.72elapsed 98%CPU (0avgtext+0avgdata 0maxresident)k')
```

Using quieve is twice as fast as Sage's general factor command in this example. Note that Sage's general factor command does nothing but call Pari's factor C library function.

```
sage: time factor(next_prime(2^90)*next_prime(2^91)) # not tested
CPU times: user 28.71 s, sys: 0.28 s, total: 28.98 s
Wall time: 29.38 s
1237940039285380274899124357 * 2475880078570760549798248507
```

Obviously, Sage's factor command should not just call Pari, but nobody has gotten around to rewriting it yet.

GMP-ECM Paul Zimmerman's GMP-ECM is included in Sage. The elliptic curve factorization (ECM) algorithm is the best algorithm for factoring numbers of the form n=pm, where p is not "too big". ECM is an algorithm due to Hendrik Lenstra, which works by "pretending" that n is prime, choosing a random elliptic curve over $\mathbf{Z}/n\mathbf{Z}$, and doing arithmetic on that curve—if something goes wrong when doing arithmetic, we factor n.

In the following example, GMP-ECM is over 10 times faster than Sage's generic factor function. Again, this emphasizes that Sage's generic factor command would benefit from a rewrite that uses GMP-ECM and qsieve.

Elliptic Curves

Cremona's Databases Cremona's databases of elliptic curves are part of Sage. The curves up to conductor 10,000 come standard with Sage, and an there is an optional download to gain access to his complete tables. From a shell, you should run

```
sage -i database_cremona_ellcurve
```

to automatically download and install the extended table.

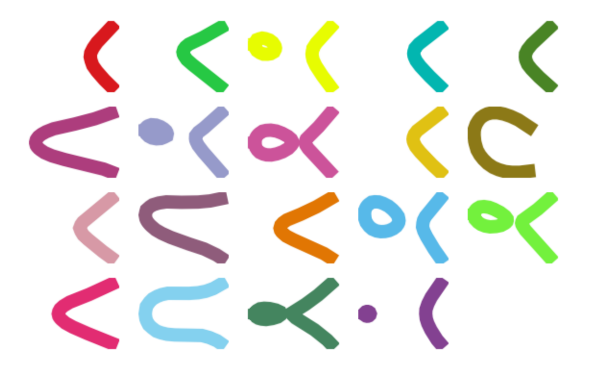
To use the database, just create a curve by giving

```
sage: EllipticCurve('5077a1')
Elliptic Curve defined by y^2 + y = x^3 - 7*x + 6 over Rational Field
sage: C = CremonaDatabase()
sage: C[37]['allcurves']
{'a1': [[0, 0, 1, -1, 0], 1, 1],
'b1': [[0, 1, 1, -23, -50], 0, 3],
'b2': [[0, 1, 1, -1873, -31833], 0, 1],
'b3': [[0, 1, 1, -3, 1], 0, 3]}
sage: C.isogeny_class('37b')
[Elliptic Curve defined by y^2 + y = x^3 + x^2 - 23*x - 50
over Rational Field, ...]
```

There is also a Stein-Watkins database that contains hundreds of millions of elliptic curves. It's over a 2GB download though!

Bryan Birch's Birthday Card Bryan Birch recently had a birthday conference, and I used Sage to draw the cover of his birthday card by enumerating all optimal elliptic curves of conductor up to 37, then plotting them with thick randomly colored lines. As you can see below, plotting an elliptic curve is as simple as calling the plot method on it. Also, the graphics array command allows us to easily combine numerous plots into a single graphics object.

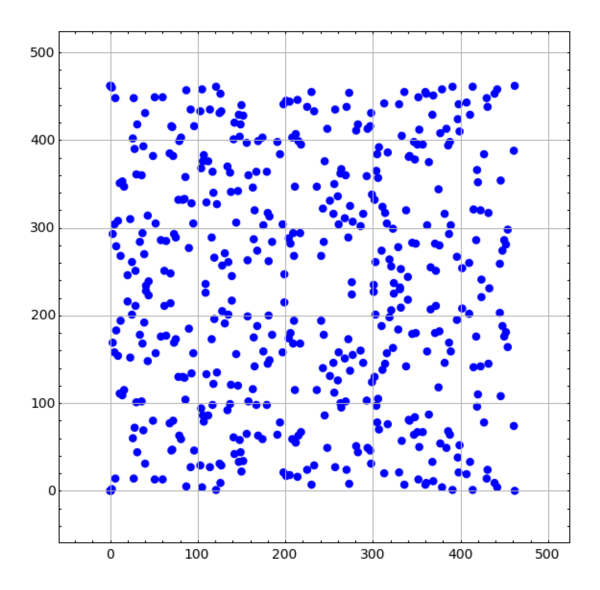
```
sage: v = cremona_optimal_curves([11..37])
sage: w = [E.plot(thickness=10,rgbcolor=(random(),random(),random())) for E in v]
sage: graphics_array(w, 4, 5).show(axes=False)
```



Plotting Modulo p We can use Sage's interact feature to draw a plot of an elliptic curve modulo p, with a slider that one drags to change the prime p. The interact feature of Sage is very helpful for interactively changing parameters and viewing the results. Type interact? for more help and examples and visit the web page http://wiki.sagemath.org/interact.

In the code below we first define the elliptic curve E using the Cremona label 37a. Then we define an interactive function f, which is made interactive using the @interact Python decorator. Because the default for p is primes(2,500), the Sage notebook constructs a slider that varies over the primes up to 500. When you drag the slider and let go, a plot is drawn of the affine \mathbf{F}_p points on the curve $E_{\mathbf{F}_p}$. Of course, one should never plot curves over finite fields, which makes this even more fun.

```
E = EllipticCurve('37a')
@interact
def f(p=primes(2,500)):
    show(plot(E.change_ring(GF(p)),pointsize=30),
    axes=False, frame=True, gridlines="automatic",
    aspect_ratio=1, gridlinesstyle={'rgbcolor':(0.7,0.7,0.7)})
```



Schoof-Elkies-Atkin Point Counting Sage includes sea.gp, which is a fast implementation of the SEA (Schoff-Elkies-Atkin) algorithm for counting the number of points on an elliptic curve over \mathbf{F}_p .

We create the finite field $k = \mathbf{F}_p$, where p is the next prime after 10^{20} . The next prime command uses Pari's nextprime function, but proves primality of the result (unlike Pari which gives only the next probable prime after a number). Sage also has a next probable prime function.

```
sage: k = GF(next\_prime(10^20))
```

compute its cardinality, which behind the scenes uses SEA.

```
sage: E = EllipticCurve_from_j(k.random_element())
sage: E.cardinality() # less than a second
9999999999971984255
```

To see how Sage chooses when to use SEA versus other methods, type E.cardinality?? and read the source code. As of this writing, it simply uses SEA whenever $p > 10^{18}$.

p-adic Regulators Sage has the world's best code for computing p-adic regulators of elliptic curves, thanks to work of David Harvey and Robert Bradshaw. The p-adic regulator of an elliptic curve E at a good ordinary prime p is the determinant of the global p-adic height pairing matrix on the Mordell-Weil group $E(\mathbf{Q})$. (This has nothing to do with local or Archimedean heights.) This is the analogue of the regulator in the Mazur-Tate-Teitelbaum p-adic analogue of the Birch and Swinnerton-Dyer conjecture.

In particular, Sage implements Harvey's improvement on an algorithm of Mazur-Stein-Tate, which builds on Kiran Kedlaya's Monsky-Washnitzer approach to computing p-adic cohomology groups.

We create the elliptic curve with Cremona label 389a, which is the curve of smallest conductor and rank 2. We then compute both the 5-adic and 997-adic regulators of this curve.

Before the new algorithm mentioned above, even computing a 7-adic regulator to 3 digits of precision was a nontrivial computational challenge. Now in Sage computing the 100003-adic regulator is routine:

```
sage: E.padic_regulator(100003,5) # a couple of seconds
42582*100003^2 + 35250*100003^3 + 12790*100003^4 + 64078*100003^5 + O(100003^6)
```

p-adic L-functions p-adic L-functions play a central role in the arithmetic study of elliptic curves. They are p-adic analogues of complex analytic L-function, and their leading coefficient (at 0) is the analogue of $L^{(r)}(E,1)/\Omega_E$ in the p-adic analogue of the Birch and Swinnerton-Dyer conjecture. They also appear in theorems of Kato, Schneider, and others that prove partial results toward p-adic BSD using Iwasawa theory.

The implementation in Sage is mainly due to work of myself, Christian Wuthrich, and Robert Pollack. We use Sage to compute the 5-adic *L*-series of the elliptic curve 389a of rank 2.

```
sage: E = EllipticCurve('389a')
sage: L = E.padic_lseries(5)
sage: L
5-adic L-series of Elliptic Curve defined
by y^2 + y = x^3 + x^2 - 2*x over Rational Field
sage: L.series(3)
0(5^5) + 0(5^2)*T + (4 + 4*5 + 0(5^2))*T^2 +
(2 + 4*5 + 0(5^2))*T^3 + (3 + 0(5^2))*T^4 + 0(T^5)
```

Bounding Shafarevich-Tate Groups Sage implements code to compute numerous explicit bounds on Shafarevich-Tate Groups of elliptic curves. This functionality is only available in Sage, and uses results Kolyvagin, Kato, Perrin-Riou, etc., and unpublished papers of Wuthrich and me.

```
sage: E = EllipticCurve('11a1')
sage: E.sha().bound()  # so only 2 could divide sha
[2]
sage: E = EllipticCurve('37a1') # so only 2 could divide sha
sage: E.sha().bound()
([2], 1)
sage: E = EllipticCurve('389a1')
sage: E.sha().bound()
(0, 0)
```

The (0,0) in the last output above indicates that the Euler systems results of Kolyvagin and Kato give no information about finiteness of the Shafarevich-Tate group of the curve E. In fact, it is an open problem to prove this finiteness, since E has rank 2, and finiteness is only known for elliptic curves for which $L(E,1) \neq 0$ or $L'(E,1) \neq 0$.

Partial results of Kato, Schneider and others on the p-adic analogue of the BSD conjecture yield algorithms for bounding the p-part of the Shafarevich-Tate group. These algorithms require as input explicit computation of p-adic L-functions, p-adic regulators, etc., as explained in Stein-Wuthrich. For example, below we use Sage to prove that 5 and 7 do not divide the Shafarevich-Tate group of our rank 2 curve 389a.

```
sage: E = EllipticCurve('389a1')
sage: sha = E.sha()
sage: sha.p_primary_bound(5)  # iwasawa theory ==> 5 doesn't divide sha
0
sage: sha.p_primary_bound(7)  # iwasawa theory ==> 7 doesn't divide sha
```

This is consistent with the Birch and Swinnerton-Dyer conjecture, which predicts that the Shafarevich-Tate group is trivial. Below we compute this predicted order, which is the floating point number 1.000000 to some precision. That the result is a floating point number helps emphasize that it is an open problem to show that the conjectural order of the Shafarevich-Tate group is even a rational number in general!

```
sage: E.sha().an()
1.000000000000000
```

Mordell-Weil Groups and Integral Points Sage includes both Cremona's mwrank library and Simon's 2-descent GP scripts for computing Mordell-Weil groups of elliptic curves.

```
sage: E = EllipticCurve([1,2,5,17,159])
sage: E.conductor() # not in the Tables
10272987
sage: E.gens() # a few seconds
[(-3 : 9 : 1), (-3347/3249 : 1873597/185193 : 1)]
```

Sage can also compute the torsion subgroup, isogeny class, determine images of Galois representations, determine reduction types, and includes a full implementation of Tate's algorithm over number fields.

Sage has the world's fastest implementation of computation of all integral points on an elliptic curve over Q, due to work of Cremona, Michael Mardaus, and Tobias Nagel. This is also the only free open source implementation available.

```
sage: E = EllipticCurve([1,2,5,7,17])
sage: E.integral_points(both_signs=True)
[(1 : -9 : 1), (1 : 3 : 1)]
```

A very impressive example is the lowest conductor elliptic curve of rank 3, which has 36 integral points.

```
sage: E = elliptic_curves.rank(3)[0]
sage: E.integral_points(both_signs=True) # less than 3 seconds
[(-3 : -1 : 1), (-3 : 0 : 1), (-2 : -4 : 1), (-2 : 3 : 1), ...(816 : -23310 : 1), (816 : 23309 : 1)]
```

The algorithm to compute all integral points involves first computing the Mordell-Weil group, then bounding the integral points, and listing all integral points satisfying those bounds. See Cohen's new GTM 239 for complete details.

The complexity grows exponentially in the rank of the curve. We can do the above calculation, but with the first known curve of rank 4, and it finishes in about a minute (and outputs 64 points).

```
sage: E = elliptic_curves.rank(4)[0]
sage: E.integral_points(both_signs=True) # about a minute
[(-10 : 3 : 1), (-10 : 7 : 1), ...
(19405 : -2712802 : 1), (19405 : 2693397 : 1)]
```

L-functions

Evaluation We next compute with the complex L-function

$$L(E,s) = \prod_{p|\Delta=389} \frac{1}{1 - a_p p^{-s} + p p^{-2s}} \cdot \prod_{p|\Delta=389} \frac{1}{1 - a_p p^{-s}}$$

of E. Though the above Euler product only defines an analytic function on the right half plane where Re(s) > 3/2, a deep theorem of Wiles et al. (the Modularity Theorem) implies that it has an analytic continuation to the whole complex plane and functional equation. We can evaluate the function L anywhere on the complex plane using Sage (via code of Tim Dokchitser).

Taylor Series We can also compute the Taylor series of L about any point, thanks to Tim Dokchitser's code.

```
sage: E = EllipticCurve('389a1')
sage: L = E.lseries()
sage: Ld = L.dokchitser()
sage: Ld.taylor_series(1,4) #random due to numerical noise
-1.28158145691931e-23 + (7.26268290635587e-24)*z + 0.759316500288427*z^2 - 0.430302337583362*z^3 + 0
```

GRH The Generalized Riemann Hypothesis asserts that all nontrivial zeros of L(E,s) are of the form 1+iy. Mike Rubinstein has written a C++ program that is part of Sage that can for any n compute the first n values of y such that 1+iy is a zero of L(E,s). It also verifies the Riemann Hypothesis for these zeros (I think). Rubinstein's program can also do similar computations for a wide class of L-functions, though not all of this functionality is as easy to use from Sage as for elliptic curves. Below we compute the first 10 zeros of L(E,s), where E is still the rank 2 curve 389a.

```
sage: L.zeros(10)
[0.000000000, 0.000000000, 2.87609907, 4.41689608, 5.79340263,
6.98596665, 7.47490750, 8.63320525, 9.63307880, 10.3514333]
```

The Matrix of Frobenius on Hyperelliptic Curves

Sage has a highly optimized implementation of the Harvey-Kedlaya algorithm for computing the matrix of Frobenius associated to a curve over a finite field. This is an implementation by David Harvey, which is GPL'd and depends only on NTL and zn_poly (a C library in Sage for fast arithmetic in $(\mathbf{Z}/n\mathbf{Z})[x]$).

We import the hypellfrob function and call it on a polynomial over **Z**.

We do the same calculation but in $\mathbb{Z}/101^4\mathbb{Z}$, which gives enough precision to recognize the exact characteristic polynomial in $\mathbb{Z}[x]$ of Frobenius as an element of the endomorphism ring. This computation is still very fast, taking only a fraction of a second.

```
sage: M = hypellfrob(p, 4, f) # about 0.25 seconds
sage: M[0,0]
91844754 + O(101^4)
```

The characteristic polynomial of Frobenius is $x^4 + 7x^3 + 167x^2 + 707x + 10201$, which determines the ζ function of the curve $y^2 = f(x)$.

```
sage: M.charpoly()
(1 + O(101^4)) *x^4 + (7 + O(101^4)) *x^3 + (167 + O(101^4)) *x^2 + (707 + O(101^4)) *x + (10201 + O(101^4))
```

Note: Exercise: Write down zeta function explicitly, count points over some finite fields and see that things match up.

Modular Symbols

Modular symbols play a key role in algorithms for computing with modular forms, special values of L-functions, elliptic curves, and modular abelian varieties. Sage has the most general implementation of modular symbols available, thanks to work of myself, Jordi Quer (of Barcelona) and Craig Citro (a student of Hida). Moreover, computation with modular symbols is by far my most favorite part of computational mathematics. There is still a lot of tuning and optimization work to be done for modular symbols in Sage, in order for it to be across the board the fastest implementation in the world, since my Magma implementation is still better in some important cases.

Note: I will talk much more about modular symbols in my lecture on Friday, which will be about modular forms and related objects.

We create the space M of weight 4 modular symbols for a certain congruence subgroup $\Gamma_H(13)$ of level 13. Then we compute a basis for this space, expressed in terms of Manin symbols. Finally, we compute the Hecke operator T_2 acting on M, find its characteristic polynomial and factor it. We also compute the dimension of the cuspidal subspace.

```
sage: M = ModularSymbols(GammaH(13,[3]), weight=4)
sage: M
Modular Symbols space of dimension 14 for Congruence Subgroup
Gamma_H(13) with H generated by [3] of weight 4 with sign 0
and over Rational Field
sage: M.basis()
([X^2,(0,1)], [X^2,(0,7)], [X^2,(2,5)], [X^2,(2,8)], [X^2,(2,9)],
[X^2,(2,10)], [X^2,(2,11)], [X^2,(2,12)], [X^2,(4,0)], [X^2,(4,3)],
[X^2,(4,6)], [X^2,(4,8)], [X^2,(4,12)], [X^2,(7,1)])
sage: factor(charpoly(M.T(2)))
```

```
(x - 7) * (x + 7) * (x - 9)^2 * (x + 5)^2 * (x^2 - x - 4)^2 * (x^2 + 9)^2

sage: dimension(M.cuspidal_subspace())
```

{Cremona's Modular Symbols Library} Sage includes John Cremona's specialized and insanely fast implementation of modular symbols for weight 2 and trivial character. We illustrate below computing the space of modular symbols of level 20014, which has dimension 5005, along with a Hecke operator on this space. The whole computation below takes only a few seconds; a similar computation takes a few minutes using Sage's generic modular symbols code. Moreover, Cremona has done computations at levels over 200,000 using his library, so the code is known to scale well to large problems. The new code in Sage for modular symbols is much more general, but doesn't scale nearly so well (yet).

```
sage: M = CremonaModularSymbols(20014) # few seconds
sage: M
Cremona Modular Symbols space of dimension 5005 for
Gamma_0(20014) of weight 2 with sign 0
sage: t = M.hecke_matrix(3) # few seconds
```

Enumerating Totally Real Number Fields

As part of his project to enumerate Shimura curves, John Voight has contributed code to Sage for enumerating totally real number fields. The algorithm isn't extremely complicated, but it involves some "inner loops" that have to be coded to run very quickly. Using Cython, Voight was able to implement exactly the variant of Newton iteration that he needed for his problem.

The function enumerate_totallyreal_fields_prim(n, B, ...) enumerates without using a database (!) primitive (no proper subfield) totally real fields of degree n > 1 with discriminant $d \le B$.

We compute the totally real quadratic fields of discriminant ≤ 50 . The calculation below, which is almost instant, is done in real time and is not a table lookup.

```
sage: enumerate_totallyreal_fields_prim(2,50) [[5, x^2 - x - 1], [8, x^2 - 2], [12, x^2 - 3], [13, x^2 - x - 3], [17, x^2 - x - 4], [21, x^2 - x - 5], [24, x^2 - 6], [28, x^2 - 7], [29, x^2 - x - 7], [33, x^2 - x - 8], [37, x^2 - x - 9], [40, x^2 - 10], [41, x^2 - x - 10], [44, x^2 - 11]]
```

We compute all totally real quintic fields of discriminant $\leq 10^5$. Again, this is done in real time - it's not a table lookup!

```
sage: enumerate_totallyreal_fields_prim(5,10^5)
[[14641, x^5 - x^4 - 4*x^3 + 3*x^2 + 3*x - 1],
[24217, x^5 - 5*x^3 - x^2 + 3*x + 1],
[36497, x^5 - 2*x^4 - 3*x^3 + 5*x^2 + x - 1],
[38569, x^5 - 5*x^3 + 4*x - 1],
[65657, x^5 - x^4 - 5*x^3 + 2*x^2 + 5*x + 1],
[70601, x^5 - x^4 - 5*x^3 + 2*x^2 + 3*x - 1],
[81509, x^5 - x^4 - 5*x^3 + 3*x^2 + 5*x - 2],
[81589, x^5 - 6*x^3 + 8*x - 1],
[89417, x^5 - 6*x^3 - x^2 + 8*x + 3]]
```

Bernoulli Numbers

Mathematica and Pari From the Mathematica website:

"Today We Broke the Bernoulli Record: From the Analytical Engine to Mathematica April 29, 2008 Oleksandr Pavlyk, Kernel Technology A week ago, I took our latest development version of Mathematica, and I typed BernoulliB[10^7]. And then I waited. Yesterday–5 days, 23 hours, 51 minutes, and 37 seconds later–I got the result!"

Tom Boothby did that same computation in Sage, which uses Pari's bernfrac command that uses evaluation of ζ and factorial to high precision, and it took 2 days, 12 hours.

David Harvey's bernmm Then David Harvey came up with an entirely new algorithm that parallelizes well. He gives these timings for computing B_{10^7} on his machine (it takes 59 minutes, 57 seconds on my 16-core 1.8ghz Opteron box):

```
PARI: 75 h, Mathematica: 142 h
bernmm (1 core) = 11.1 h, bernmm (10 cores) = 1.3 h
```

"Running on 10 cores for 5.5 days, I [David Harvey] computed [the Bernoulli number] B_k for $k = 10^8$, which I believe is a new record. Essentially it's the multimodular algorithm I suggested earlier on this thread, but I figured out some tricks to optimise the crap out of the computation of $B_k \mod p$."

So now Sage is the fastest in the world for large Bernoulli numbers. The timings below are on a 24-core 2.66Ghz Xeon box.

```
sage: w1 = bernoulli(100000, num_threads=16) # 0.9 seconds wall time
sage: w2 = bernoulli(100000, algorithm='pari') # long time (6s on sage.math, 2011)
sage: w1 == w2 # long time
True
```

Polynomial Arithmetic

FLINT: Univariate Polynomial Arithmetic Sage uses Bill Hart and David Harvey's GPL'd Flint C library for arithmetic in $\mathbb{Z}[x]$. Its main claim to fame is that it is the world's fastest for polynomial multiplication, e.g., in the benchmark below it is faster than NTL and Magma on some systems (though such benchmarks of course change as software improves). Behind the scenes Flint contains some carefully tuned discrete Fourier transform code.

```
sage: Rflint = PolynomialRing(ZZ, 'x')
sage: f = Rflint([ZZ.random_element(2^64) for _ in [1..32]])
sage: g = Rflint([ZZ.random_element(2^64) for _ in [1..32]])
sage: timeit('f*q')
                                  # random output
625 loops, best of 3: 105 microseconds per loop
sage: Rntl = PolynomialRing(ZZ, 'x', implementation='NTL')
sage: f = Rntl([ZZ.random_element(2^64) for _ in [1..32]])
sage: g = Rntl([ZZ.random_element(2^64) for _ in [1..32]])
sage: timeit('f*q')
                                  # random output
625 loops, best of 3: 310 microseconds per loop
sage: ff = magma(f); qq = magma(q) #optional - magma
sage: s = 'time \ v := [\$s * \$s : i in [1..10^5]];'\$(ff.name(), gg.name()) #optional - magma
sage: magma.eval(s) #optional - magma
'Time: ...'
```

Singular: Multivariate Polynomial Arithmetic Multivariate polynomial arithmetic in many cases uses Singular in library mode (due to Martin Albrecht), which is quite fast. For example, below we do the Fateman benchmark over the finite field of order 32003, and compare the timing with Magma.

```
sage: P.<x,y,z> = GF(32003)[]
sage: p = (x+y+z+1)^10
sage: q = p+1
sage: timeit('p*q')  # random output
125 loops, best of 3: 1.53 ms per loop

sage: p = (x+y+z+1)^20
sage: q = p+1
sage: timeit('p*q')  # not tested - timeout if SAGE_DEBUG=yes
5 loops, best of 3: 384 ms per loop

sage: pp = magma(p); qq = magma(q) #optional - magma
sage: pp = magma(p); qq = magma(q) #optional - magma
sage: s = 'time w := %s*%s;'%(pp.name(),qq.name()) #optional - magma
sage: magma.eval(s) #optional - magma
'Time: ...'
```

Modular Forms

This section is about computing with modular forms, modular symbols, and modular abelian varieties. Most of the Sage functionality we describe below is new code written for Sage by myself, Craig Citro, Robert Bradshaw, and Jordi Quer in consultation with John Cremona. It has much overlap in functionality with the modular forms code in Magma, which I developed during 1998-2004.

Modular Forms and Hecke Operators

Congruence subgroups

Definition A congruence subgroup is a subgroup of the group $SL_2(\mathbf{Z})$ of determinant ± 1 integer matrices that contains

$$\Gamma(N) = \operatorname{Ker}(\operatorname{SL}_2(\mathbf{Z}) \to \operatorname{SL}_2(\mathbf{Z}/N\mathbf{Z}))$$

for some positive integer N. Since $\Gamma(N)$ has finite index in $SL_2(\mathbf{Z})$, all congruence subgroups have finite index. The converse is not true, though in many other settings it is true (see [paper of Serre]).

The inverse image $\Gamma_0(N)$ of the subgroup of upper triangular matrices in $\operatorname{SL}_2(\mathbf{Z}/N\mathbf{Z})$ is a congruence subgroup, as is the inverse image $\Gamma_1(N)$ of the subgroup of matrices of the form $\begin{pmatrix} 1 & * \\ 0 & 1 \end{pmatrix}$. Also, for any subgroup $H \subset (\mathbf{Z}/N\mathbf{Z})^*$, the inverse image $\Gamma_H(N)$ of the subgroup of $\operatorname{SL}_2(\mathbf{Z}/N\mathbf{Z})$ of all elements of the form $\begin{pmatrix} a & * \\ 0 & d \end{pmatrix}$ with $d \in H$ is a congruence subgroup.

We can create each of the above congruence subgroups in Sage, using the Gamma0, Gamma1, and GammaH commands.

```
sage: Gamma0(8)
Congruence Subgroup Gamma0(8)
sage: Gamma1(13)
Congruence Subgroup Gamma1(13)
sage: GammaH(11,[4])
Congruence Subgroup Gamma_H(11) with H generated by [4]
```

The second argument to the GammaH command is a list of generators of the subgroup H of $(\mathbf{Z}/N\mathbf{Z})^*$.

Generators Sage can compute a list of generators for these subgroups. The algorithm Sage uses is a straightforward generic procedure that uses coset representatives for the congruence subgroup (which are easy to enumerate) to obtain a list of generators [[ref my modular forms book]].

The list of generators Sage computes is unfortunately large. Improving this would be an excellent Sage development project, which would involve much beautiful mathematics.

UPDATE (March 2012): The project referred to above has been carried out (by several people, notably Hartmut Monien, building on earlier work of Chris Kurth). Sage now uses a much more advanced algorithm based on Farey symbols which calculates a *minimal* set of generators.

```
sage: Gamma0(2).gens()
(
[1 1] [ 1 -1]
[0 1], [ 2 -1]
)
sage: Gamma0(2).gens(algorithm="todd-coxeter") # the old implementation
(
[1 1] [-1 0] [ 1 -1] [ 1 -1] [-1 1]
[0 1], [ 0 -1], [ 0 1], [ 2 -1], [-2 1]
)
sage: len(Gamma1(13).gens())
```

Modular Forms

Definition A modular form on a congruence subgroup Γ of integer weight k is a holomorphic function f(z) on the upper half plane

$$\mathfrak{h}^* = \{ z \in \mathbf{C} : \Im(z) > 0 \} \cup \mathbf{Q} \cup \{ i \infty \}$$

such that for every matrix $\begin{pmatrix} a & b \\ c & d \end{pmatrix} \in \Gamma$, we have

$$f\left(\frac{az+b}{cz+d}\right) = (cz+d)^k f(z).$$

A cusp form is a modular form that vanishes at all of the cusps $\mathbf{Q} \cup \{i\infty\}$.

If Γ contains $\Gamma_1(N)$ for some N, then $\begin{pmatrix} 1 & 1 \\ 0 & 1 \end{pmatrix} \in \Gamma$, so the modular form condition implies that f(z) = f(z+1). This, coupled with the holomorphicity condition, implies that f(z) has a Fourier expansion

$$f(z) = \sum_{n=0}^{\infty} a_n e^{2\pi i n z}$$

with $a_n \in \mathbb{C}$. We let $q = e^{2\pi i z}$, and call $f = \sum_{n=0}^{\infty} a_n q^n$ the q-expansion of f.

Creation in Sage Henceforth we assume that Γ is either $\Gamma_1(N)$, $\Gamma_0(N)$, or $\Gamma_H(N)$ for some H and N. The complex vector space $M_k(\Gamma)$ of all modular forms of weight k on Γ is a finite dimensional vector space.

We create the space $M_k(\Gamma)$ in Sage by typing ModularForms (G, k) where G is the congruence subgroup and k is the weight.

```
sage: ModularForms(Gamma0(25), 4)
Modular Forms space of dimension 11 for ...
sage: S = CuspForms(Gamma0(25), 4, prec=15); S
Cuspidal subspace of dimension 5 of Modular Forms space ...
```

```
sage: S.basis()
[
q + q^9 - 8*q^11 - 8*q^14 + O(q^15),
q^2 - q^7 - q^8 - 7*q^12 + 7*q^13 + O(q^15),
q^3 + q^7 - 2*q^8 - 6*q^12 - 5*q^13 + O(q^15),
q^4 - q^6 - 3*q^9 + 5*q^11 - 2*q^14 + O(q^15),
q^5 - 4*q^10 + O(q^15)
]
```

Dimension Formulas Sage computes the dimensions of all these spaces using simple arithmetic formulas instead of actually computing bases for the spaces in question. In fact, Sage has the most general collection of modular forms dimension formulas of any software; type help(sage.modular.dims) to see a list of arithmetic functions that are used to implement these dimension formulas.

```
sage: ModularForms(Gamma1(949284), 456).dimension()
11156973844800
sage: a = [dimension_cusp_forms(Gamma0(N),2) for N in [1..25]]; a
[0, 0, ..., 1, 0, 0, 1, 1, 0, 1, 1, 1, 2, 2, 1, 0]
sage: oeis(a)  # optional - internet
0: A001617: Genus of modular group GAMMA_0 (n). Or, genus of modular curve X_0(n).
```

Sage doesn't have simple formulas for dimensions of spaces of modular forms of weight 1, since such formulas perhaps do not exist.

Diamond Bracket Operators The space $M_k(\Gamma_1(N))$ is equipped with an action of $(\mathbf{Z}/N\mathbf{Z})^*$ by diamond bracket operators $\langle d \rangle$, and this induces a decomposition

$$M_k(\Gamma_1(N)) = \bigoplus_{\varepsilon: (\mathbf{Z}/N\mathbf{Z})^* \to \mathbf{C}^*} M_k(N, \varepsilon),$$

where the sum is over all complex characters of the finite abelian group $(\mathbf{Z}/N\mathbf{Z})^*$. These characters are called Dirichlet characters, which are central in number theory.

The factors $M_k(N, \varepsilon)$ then have bases whose q-expansions are elements of R[[q]], where $R = \mathbf{Z}[\varepsilon]$ is the ring generated over \mathbf{Z} by the image of ε . We illustrate this with N = k = 5 below, where DirichletGroup will be described later.

```
sage: CuspForms(DirichletGroup(5).0, 5).basis()
[
q + (-zeta4 - 1)*q^2 + (6*zeta4 - 6)*q^3 - ... + O(q^6)
]
```

Dirichlet Characters Use the command DirichletGroup(N,R) to create the group of all Dirichlet characters of modulus N taking values in the ring R. If R is omitted, it defaults to a cyclotomic field.

```
sage: G = DirichletGroup(8); G
Group of Dirichlet characters of modulus 8 over Cyclotomic
Field of order 2 and degree 1
sage: v = G.list(); v
[Dirichlet character modulo 8 of conductor 1 mapping 7 |--> 1, 5 |--> 1,
Dirichlet character modulo 8 of conductor 4 mapping 7 |--> -1, 5 |--> 1,
Dirichlet character modulo 8 of conductor 8 mapping 7 |--> 1, 5 |--> -1,
Dirichlet character modulo 8 of conductor 8 mapping 7 |--> -1, 5 |--> -1]
sage: eps = G.0; eps
Dirichlet character modulo 8 of conductor 4 mapping 7 |--> -1, 5 |--> 1
```

```
sage: eps.values()
[0, 1, 0, -1, 0, 1, 0, -1]
```

Sage both represents Dirichlet characters by giving a "matrix", i.e., the list of images of canonical generators of $(\mathbf{Z}/N\mathbf{Z})^*$, and as vectors modulo and integer n. For years, I was torn between these two representations, until J. Quer and I realized that the best approach is to use both and make it easy to convert between them.

```
sage: parent(eps.element())
Vector space of dimension 2 over Ring of integers modulo 2
```

Given a Dirichlet character, Sage also lets you compute the associated Jacobi and Gauss sums, generalized Bernoulli numbers, the conductor, Galois orbit, etc.

Decomposing $M_k(\Gamma_1(N))$ Recall that Dirichlet characters give a decomposition

$$M_k(\Gamma_1(N)) = \bigoplus_{\varepsilon: (\mathbf{Z}/N\mathbf{Z})^* \to \mathbf{C}^*} M_k(N, \varepsilon).$$

Given a Dirichlet character ε we type ModularForms(eps, weight) to create the space of modular forms with that character and a given integer weight. For example, we create the space of forms of weight 5 with the character modulo 8 above that is -1 on 3 and 1 on 5 as follows.

```
sage: ModularForms(eps,5)
Modular Forms space of dimension 6, character [-1, 1] and
weight 5 over Rational Field
sage: sum([ModularForms(eps,5).dimension() for eps in v])
11
sage: ModularForms(Gamma1(8),5)
Modular Forms space of dimension 11 ...
```

Note: Exercise: Compute the dimensions of all spaces $M_2(37, \varepsilon)$ for all Dirichlet characters ε .

Hecke Operators The space $M_k(\Gamma)$ is equipped with an action of a commuting ring \mathbb{T} of Hecke operators T_n for $n \geq 1$. A standard computational problem in the theory of modular forms is to compute an explicit basis of q-expansion for $M_k(\Gamma)$ along with matrices for the action of any Hecke operator T_n , and to compute the subspace $S_k(\Gamma)$ of cusp forms.

```
sage: M = ModularForms(Gamma0(11),4)
sage: M.basis()
[
q + 3*q^3 - 6*q^4 - 7*q^5 + O(q^6),
q^2 - 4*q^3 + 2*q^4 + 8*q^5 + O(q^6),
1 + O(q^6),
q + 9*q^2 + 28*q^3 + 73*q^4 + 126*q^5 + O(q^6)
]
sage: M.hecke_matrix(2)
[0 2 0 0]
[1 2 0 0]
[0 0 9 0]
[0 0 9 0]
```

We can also compute Hecke operators on the cuspidal subspace.

```
sage: S = M.cuspidal_subspace()
sage: S.hecke_matrix(2)
[0 2]
[1 2]
sage: S.hecke_matrix(3)
[ 3 -8]
[-4 -5]
```

Hecke Operator on $M_k(\Gamma_1(N))$ At the time these lectures were first written, Sage didn't yet implement computation of the Hecke operators on $M_k(\Gamma_1(N))$, but these have subsequently been added:

These are calculated by first calculating Hecke operators on modular symbols for $\Gamma_1(N)$, which is a \mathbb{T} -module that is isomorphic to $M_k(\Gamma_1(N))$ (see *Modular Symbols*).

Modular Symbols

Modular symbols are a beautiful piece of mathematics that was developed since the 1960s by Birch, Manin, Shokorov, Mazur, Merel, Cremona, and others. Not only are modular symbols a powerful computational tool as we will see, they have also been used to prove rationality results for special values of *L*-series, to construct *p*-adic *L*-series, and they play a key role in Merel's proof of the uniform boundedness theorem for torsion points on elliptic curves over number fields

We view modular symbols as a remarkably flexible computational tool that provides a single uniform algorithm for computing $M_k(N,\varepsilon)$ for any N,ε and $k\geq 2$. There are ways to use computation of those spaces to obtain explicit basis for spaces of weight 1 and half-integral weight, so in a sense modular symbols yield everything. There are also generalizations of modular symbols to higher rank groups, though Sage currently has no code for modular symbols on higher rank groups.

Definition A modular symbol of weight k, and level N, with character ε is a sum of terms $X^iY^{k-2-i}\{\alpha,\beta\}$, where $0 \le i \le k-2$ and $\alpha,\beta \in \mathbb{P}^1(\mathbf{Q}) = \mathbf{Q} \cup \{\infty\}$. Modular symbols satisfy the relations

$$X^iY^{k-2-i}\{\alpha,\beta\}+X^iY^{k-2-i}\{\beta,\gamma\}+X^iY^{k-2-i}\{\gamma,\alpha\}=0$$

$$X^{i}Y^{k-2-i}\{\alpha,\beta\} = -X^{i}Y^{k-2-i}\{\beta,\alpha\},$$

and for every $\gamma = \begin{pmatrix} a & b \\ c & d \end{pmatrix} \in \Gamma_0(N)$, we have

$$(dX - bY)^i(-cX + aY)^{k-2-i}\{\gamma(\alpha), \gamma(\beta)\} = \varepsilon(d)X^iY^{k-2-i}\{\alpha, \beta\}.$$

The modular symbols space $\mathcal{M}_k(N,\varepsilon)$ is the torsion free $\mathbf{Q}[\varepsilon]$ -module generated by all sums of modular symbols, modulo the relations listed above. Here $\mathbf{Q}[\varepsilon]$ is the ring generated by the values of the character ε , so it is of the form $\mathbf{Q}[\zeta_m]$ for some integer m.

The amazing theorem that makes modular symbols useful is that there is an explicit description of an action of a Hecke algebra \mathbb{T} on $\mathcal{M}_k(N,\varepsilon)$, and there is an isomorphism

$$\mathcal{M}_k(N,\varepsilon;\mathbf{C}) \xrightarrow{\approx} M_k(N,\varepsilon) \oplus S_k(N,\varepsilon).$$

This means that if modular symbols are computable (they are!), then they can be used to compute a lot about the \mathbb{T} -module $M_k(N,\varepsilon)$.

Manin Symbols

Definition Though $\mathcal{M}_k(N,\varepsilon)$ as described above is not explicitly generated by finitely many elements, it is finitely generated. Manin, Shokoruv, and Merel give an explicit description of finitely many generators (Manin symbols) for this space, along with all explicit relations that these generators satisfy (see my book). In particular, if we let

$$(i, c, d) = [X^{i}Y^{2-k-i}, (c, d)] = (dX - bY)^{i}(-cX + aY)^{k-2-i}\{\gamma(0), \gamma(\infty)\},\$$

where $\gamma = \begin{pmatrix} a & b \\ c & d \end{pmatrix}$, then the Manin symbols (i, c, d) with $0 \le i \le k - 2$ and $(c, d) \in \mathbb{P}^1(N)$ generate $\mathcal{M}_k(N, \varepsilon)$.

Computing in Sage We compute a basis for the space of weight 4 modular symbols for $\Gamma_0(11)$, then coerce in (2,0,1) and (1,1,3).

We compute a modular symbols representation for the Manin symbol (2, 1, 6), and verify this by converting back.

```
sage: a = M.1; a
[X^2,(1,6)]
sage: a.modular_symbol_rep()
36*X^2*{-1/6, 0} + 12*X*Y*{-1/6, 0} + Y^2*{-1/6, 0}
sage: 36*M([2,-1/6,0]) + 12*M([1,-1/6,0]) + M([0,-1/6,0])
[X^2,(1,6)]
```

Method of Graphs

The Mestre Method of Graphs is an intriguing algorithm for computing the action of Hecke operators on yet another module X that is isomorphic to $M_2(\Gamma_0(N))$. The implementation in Sage unfortunately only works when N is prime; in contrast, my implementation in Magma works when N=pM and $S_2(\Gamma_0(M))=0$.

The matrices of Hecke operators on X are vastly sparser than on any basis of $M_2(\Gamma_0(N))$ that you are likely to use.

The method of graphs is also used in computer science to construct expander graphs with good properties. And it is important in my algorithm for computing Tamagawa numbers of purely toric modular abelian varieties. This algorithm is not implemented in Sage yet, since it is only interesting in the case of non-prime level, as it turns out.

Level One Modular Forms

Computing Δ The modular form

$$\Delta = q \prod (1 - q^n)^{24} = \sum \tau(n)q^n$$

is perhaps the world's most famous modular form. We compute some terms from the definition.

```
sage: R.\langle q \rangle = QQ[[]]

sage: q * prod( 1-q^n+0(q^6) for n in (1..5) )^24

q - 24*q^2 + 252*q^3 - 1472*q^4 + 4830*q^5 - 6048*q^6 + O(q^7)
```

There are much better ways to compute Δ , which amount to just a few polynomial multiplications over **Z**.

```
sage: D = delta_qexp(10^5)  # less than 10 seconds
sage: D[:10]
q - 24*q^2 + 252*q^3 - 1472*q^4 + ...
sage: [p for p in primes(10^5) if D[p] % p == 0]
[2, 3, 5, 7, 2411]
sage: D[2411]
4542041100095889012
sage: f = eisenstein_series_qexp(12,6) - D[:6]; f
691/65520 + 2073*q^2 + 176896*q^3 + 4197825*q^4 + 48823296*q^5 + O(q^6)
sage: f % 691
O(q^6)
```

The Victor Miller Basis The Victor Miller basis for $M_k(\operatorname{SL}_2(\mathbf{Z}))$ is the reduced row echelon basis. It's a lemma that it has all integer coefficients, and a rather nice diagonal shape.

```
sage: victor_miller_basis(24, 6)
[
1 + 52416000*q^3 + 39007332000*q^4 + 6609020221440*q^5 + O(q^6),
q + 195660*q^3 + 12080128*q^4 + 44656110*q^5 + O(q^6),
q^2 - 48*q^3 + 1080*q^4 - 15040*q^5 + O(q^6)
]
sage: dimension_modular_forms(1,200)
17
sage: B = victor_miller_basis(200, 18) #5 seconds
sage: B
```

```
[
1 + 79288314420681734048660707200000*q^17 + O(q^18),
q + 2687602718106772837928968846869*q^17 + O(q^18),
...
q^16 + 96*q^17 + O(q^18)
]
```

Note: Craig Citro has made the above computation an order of magnitude faster in code he hasn't quite got into Sage yet.

"I'll clean those up and submit them soon, since I need them for something I'm working on ... I'm currently in the process of making spaces of modular forms of level one subclass the existing code, and actually take advantage of all our fast E_k and Δ computation code, as well as cleaning things up a bit."

Half Integral Weight Forms

Basmaji's Algorithm Basmaji (page 55 of his Essen thesis, "Ein Algorithmus zur Berechnung von Hecke-Operatoren und Anwendungen auf modulare Kurven", http://wstein.org/scans/papers/basmaji/).

Let $S = S_{k+1}(\varepsilon)$ be the space of cusp forms of even integer weight k+1 and character $\varepsilon = \chi \psi^{(k+1)/2}$, where ψ is the nontrivial mod-4 Dirichlet character. Let U be the subspace of $S \times S$ of elements (a,b) such that $\Theta_2 a = \Theta_3 b$. Then U is isomorphic to $S_{k/2}(\chi)$ via the map $(a,b) \mapsto a/\Theta_3$.

This algorithm is implemented in Sage. I'm sure it could be implemented in a way that is much faster than the current implementation...

```
sage: half_integral_weight_modform_basis(DirichletGroup(16,QQ).1, 3, 10)
[]
sage: half_integral_weight_modform_basis(DirichletGroup(16,QQ).1, 5, 10)
[q - 2*q^3 - 2*q^5 + 4*q^7 - q^9 + O(q^10)]
sage: half_integral_weight_modform_basis(DirichletGroup(16*7).0^2,3,30)
[q - 2*q^2 - q^9 + 2*q^14 + 6*q^18 - 2*q^21 - 4*q^22 - q^25 + O(q^30), q^2 - q^14 - 3*q^18 + 2*q^22 + O(q^30), q^4 - q^8 - q^16 + q^28 + O(q^30), q^7 - 2*q^15 + O(q^30)]
```

Generators for Rings of Modular Forms

Computing Generators For any congruence subgroup Γ , the direct sum

$$M(\Gamma) = \bigoplus_{k \ge 0} M_k(\Gamma)$$

is a ring, since the product of modular forms $f \in M_k(\Gamma)$ and $g \in M_{k'}(\Gamma)$ is an element $fg \in M_{k+k'}(\Gamma)$. Sage can compute likely generators for rings of modular forms, but currently doesn't prove any of these results.

We verify the statement proved in Serre's "A Course in Arithmetic" that E_4 and E_6 generate the space of level one modular forms.

```
sage: ModularFormsRing(SL2Z).generators(prec=4)
[(4, 1 + 240*q + 2160*q^2 + 6720*q^3 + O(q^4)),
  (6, 1 - 504*q - 16632*q^2 - 122976*q^3 + O(q^4))]
```

Have you ever wondered which forms generate the ring $M(\Gamma_0(2))$? It turns out that one form of weight 2 and one form of weight 4 suffice.

```
sage: ModularFormsRing(Gamma0(2)).generators(prec=12)
[(2, 1 + 24*q + 24*q^2 + 96*q^3 + 24*q^4 + 144*q^5 + 96*q^6 + 192*q^7 + 24*q^8 + 312*q^9 + 144*q^10
(4, 1 + 240*q^2 + 2160*q^4 + 6720*q^6 + 17520*q^8 + 30240*q^10 + O(q^12))]
```

Here's generators for $M(\Gamma_0(3))$. Notice that elements of weight 6 are now required, in addition to weights 2 and 4.

```
sage: ModularFormsRing(Gamma0(3)).generators()
[(2, 1 + 12*q + 36*q^2 + 12*q^3 + 84*q^4 + 72*q^5 + 36*q^6 + 96*q^7 + 180*q^8 + 12*q^9 + O(q^10)),
(4, 1 + 240*q^3 + 2160*q^6 + 6720*q^9 + O(q^10)),
(6, 1 - 504*q^3 - 16632*q^6 - 122976*q^9 + O(q^10))]
```

(*Note*: As of 2012, updates to the code mean that the output of this test is not quite the same as it was in 2008, but of course there are multiple equally valid answers.)

We can also handle rings of modular forms for odd congruence subgroups, but with the usual caveat that we can't calculate forms of weight 1. So these are elements generating the graded ring of forms of weight 0 or \geq 2.

```
sage: ModularFormsRing(Gamma1(3)).generators() [(2, 1 + 12*q + 36*q^2 + 12*q^3 + 84*q^4 + 72*q^5 + 36*q^6 + 96*q^7 + 180*q^8 + 12*q^9 + O(q^10)), (3, 1 + 54*q^2 + 72*q^3 + 432*q^5 + 270*q^6 + 918*q^8 + 720*q^9 + O(q^10)), (3, q + 3*q^2 + 9*q^3 + 13*q^4 + 24*q^5 + 27*q^6 + 50*q^7 + 51*q^8 + 81*q^9 + O(q^10)), (4, 1 + 240*q^3 + 2160*q^6 + 6720*q^9 + O(q^10))]
```

L-series

L-series of Δ Thanks to wrapping work of Jennifer Balakrishnan of M.I.T., we can compute explicitly with the L-series of the modular form Δ . Like for elliptic curves, behind these scenes this uses Dokchitsers L-functions calculation Pari program.

```
sage: L = delta_lseries(); L
L-series associated to the modular form Delta
sage: L(1)
0.0374412812685155
```

L-series of a Cusp Form In some cases we can also compute with L-series attached to a cusp form.

```
sage: f = CuspForms(2,8).newforms()[0]
sage: L = f.lseries()
sage: L(1)
0.0884317737041015
sage: L(0.5)
0.0296568512531983
```

L-series of a General Newform is Not Implemented Unfortunately, computing with the L-series of a general newform is not yet implemented.

```
sage: S = CuspForms(23,2); S
Cuspidal subspace of dimension 2 of Modular Forms space of
dimension 3 for Congruence Subgroup Gamma0(23) of weight
2 over Rational Field
sage: f = S.newforms('a')[0]; f
q + a0*q^2 + (-2*a0 - 1)*q^3 + (-a0 - 1)*q^4 + 2*a0*q^5 + O(q^6)
```

Computing with L(f, s) totally not implemented yet, though should be easy via Dokchitser.

Modular Abelian Varieties

The quotient of the extended upper half plane \mathfrak{h}^* by the congruence subgroup $\Gamma_1(N)$ is the modular curve $X_1(N)$. Its Jacobian $J_1(N)$ is an abelian variety that is canonically defined over \mathbf{Q} . Likewise, one defines a modular abelian variety $J_0(N)$ associated to $\Gamma_0(N)$.

A modular abelian variety is an abelian variety over \mathbf{Q} that is a quotient of $J_1(N)$ for some N.

The biggest recent theorem in number theory is the proof of Serre's conjecture by Khare and Wintenberger. According to an argument of Ribet and Serre, this implies the following modularity theorem, which generalizes the modularity theorem that Taylor-Wiles proved in the course of proving Fermat's Last Theorem.

One of my long-term research goals is to develop a systematic theory for computing with modular abelian varieties. A good start is the observation using the Abel-Jacobi theorem that every modular abelian variety (up to isomorphism) can be specified by giving a lattice in a space of modular symbols.

Computing in Sage We define some modular abelian varieties of level 39, and compute some basic invariants.

```
sage: D = J0(39).decomposition(); D
Simple abelian subvariety 39a(1,39) of dimension 1 of JO(39),
Simple abelian subvariety 39b(1,39) of dimension 2 of J0(39)
sage: D[1].lattice()
Free module of degree 6 and rank 4 over Integer Ring
Echelon basis matrix:
[ 1 0 0 1 -1 0]
       1 0 -1
    1
                0.1
[ 0 0 2 0 -1 0]
[ 0 0 0 0 0 1]
sage: G = D[1].rational_torsion_subgroup(); G
Torsion subgroup of Simple abelian subvariety 39b(1,39)
of dimension 2 of J0(39)
sage: G.order()
sage: G.gens()
[[(1/14, 2/7, 0, 1/14, -3/14, 1/7)], [(0, 1, 0, 0, -1/2, 0)],
[(0, 0, 1, 0, -1/2, 0)]]
sage: B, phi = D[1]/G
sage: B
Abelian variety factor of dimension 2 of J0(39)
sage: phi.kernel()
(Finite subgroup with invariants [2, 14] ...
```

Endomorphisms There is an algorithm in Sage for computing the exact endomorphism ring of any modular abelian variety.

```
sage: A = J0(91)[2]; A
Simple abelian subvariety 91c(1,91) of dimension 2 of J0(91)
sage: R = End(A); R
Endomorphism ring of Simple abelian subvariety 91c(1,91)
of dimension 2 of J0(91)
sage: for x in R.gens(): print x.matrix(),'\n'
[1 0 0 0]
[0 1 0 0]
[0 0 1 0]
[0 0 0 1]
```

```
[ 0 4 -2 0]
[-1 5 -2 1]
[-1 2 0 2]
[-1 1 0 3]
```

It is also possible to test isomorphism of two modular abelian varieties. But much exciting theoretical and computational work remains to be done.

12.1.19 Creating a Tutorial from a Worksheet

Sage has a number of thematic tutorials and contains everything needed to turn a worksheet created in the Sage notebook (sagenb) into a tutorial.

• Once you have created a worksheet and are satisfied with the text and computations, download it to a directory.

We will assume here that the worksheet is called Tutorial.sws and the directory is called make_tutorial. We also assume that sage is your Sage command; if it is not in your PATH then replace this with the path to your Sage installation, such as /Applications/Sage-6.2.app/Contents/Resources/sage/sage if you are using the Mac app and have placed it in your Applications directory.

• Next, you will need an optional package to parse your worksheet. Use the command:

```
sage --pip install beautifulsoup4
```

to install it (or, in the Mac app, use the Terminal Session advanced menu with --pip install beautifulsoup4).

• Then we will use the sws2rst script to turn the worksheet into a document in the ReStructuredText format. Be sure you are in the same directory as the worksheet:

```
sage --sws2rst Tutorial.sws
```

This will create an .rst file along with a subdirectory of image files (which may be empty if there are no images).

You can find help for sws2rst with the command sage --sws2rst -h once you have installed beautifulsoup.

- In principle, such a file could be added directly to Sage's documentation (see the developer's manual). However, you probably want to check whether it looks right first. So next we will compile this file to html documentation.
 - Follow the instructions of sage --sws2rst --sphinxify. First, we will open a Sage shell session, where all appropriate Sage references already work properly:

```
sage --sh
```

From here, you should be able to just type:

```
sphinx-quickstart
```

and then respond to prompts for turning your .rst file into documentation. For most of them you can just hit enter/return to accept the defaults. However, you will probably want to

- * Enter a name for the project
- * Enter a name for you
- * Type y for the question about using MathJax

Keep note of the instructions; the main other thing to do is add your file's name to index.rst, and then just do:

```
make html
```

and wait while magic happens. To see the results, open the file make_tutorial/_build/html/Tutorial.html with a browser, or use your graphical file system to navigate to the same place.

• Now you can modify the .rst file more and repeat the steps of compiling it until it is ready for inclusion, or just for distribution among other Sage users as an HTML file. (Do make pdf for a PDF version.)

12.1.20 Profiling in Sage

This page lists several methods available in Sage to measure and analyze the performances of a piece of code. For more general information on profiling, see Wikipedia article Profiling_(computer_programming).

Table of contents

- Profiling in Sage
 - How long does it take? %time and %timeit
 - Python-level function calls: %prun
 - Python-level line-by-line profiling: %lprun
 - C-level function calls: %crun
 - C-level line-by-line profiling: perf (Linux only)

How long does it take? %time and %timeit

The two IPython magics %time and %timeit measure the time it takes to run a command:

```
sage: %time p=random_prime(2**300)
CPU times: user 152 ms, sys: 0 ns, total: 152 ms
Wall time: 150 ms
sage: %timeit p=random_prime(2**300)
10 loops, best of 3: 62.2 ms per loop
```

Note that while %time only runs the command once, %timeit tries to return a more meaningful value over several runs.

For more information see %timeit? or this page.

Note that Sage provides a timeit function which also runs in the Sage notebook.

Python-level function calls: %prun

With %prun, you can obtain the list of all Python functions involved in a computation, as well as the time spent on each of them:

```
sage: %prun _=random_prime(2**500)
    468 function calls in 0.439 seconds
Ordered by: internal time
```

```
ncalls tottime percall cumtime percall filename: lineno(function)
   32
       0.438
               0.014
                       0.438 0.014 {method 'is_prime' of 'sage.rings.integer.Integer' object:
                       0.439 0.014 arith.py:407(is_prime)
   32
        0.001 0.000
   32
        0.000
                 0.000
                         0.001
                                 0.000 random.py:175(randrange)
   32
        0.000
                0.000
                         0.000
                                 0.000 random.py:244(_randbelow)
```

The most time-consuming functions should appear on the top. A description of the different columns is available here.

Note: You may want to sort this list differently, e.g. use %prun -s cumulative for decreasing cumulative time.

Alternatively, you can "save" this data to a Stats object for further inspection:

```
sage: %prun -r random_prime(2**500)
sage: stats_object = _
sage: stats_object.total_calls
2547
```

For more information see %prun? or this page.

Visualize the statistics: you can obtain a more graphical output with RunSnake and Sage's function runsnake():

```
sage: runsnake('random_prime(2**500)')
```

Python-level line-by-line profiling: %lprun

With line_profiler and its %lprun magic, you can find out which lines of one (or many) functions are the most time-consuming. The syntax is the following:

```
%lprun -f function1 -f function2 code_to_run
```

This will display the line-by-line analysis of function1 and function2 when code_to_run is executed:

```
sage: %lprun -f random_prime random_prime(2**500)
Line #
         Hits
                     Time Per Hit % Time Line Contents
                                                    def random_prime(n, proof=None, lbound=2):
1193
. . .
                                                         # since we don't want current_randstate to get
1251
                                                        # pulled when you say "from sage.arith import *".
1252
                                            0.0 from sage.misc.randstate import current_randstate
0.0 from sage.structure.proof.proof import get_flag
0.0 proof = get_flag(proof, "arithmetic")
             1
                          11
                                  11.0
                                                       from sage.misc.randstate import current_randstate
1253
             1
                                   7.0
1255
             1
                           6
                                   6.0
1256
             1
                          17
                                   17.0
                                             0.0
                                                        n = ZZ(n)
```

In order to install line_profiler you must first run the following command:

```
[user@localhost ~] sage -pip install "line_profiler"
```

C-level function calls: %crun

With %crun, you can obtain the list of all C functions involved in a computation, as well as the time spent on each of them. You will need to have the Google performance analysis tools installed on your system:

```
sage: %crun p=random_prime(2**500)
PROFILE: interrupts/evictions/bytes = 45/0/18344
Total: 45 samples
                           35 77.8% PyEval_EvalCode
      0
         0.0%
                 0.0%
                            35 77.8% PyEval_EvalCodeEx
      0
          0.0%
                 0.0%
               0.0%
0.0%
0.0%
                            35 77.8% PyEval_EvalFrameEx
          0.0%
                            35 77.8% PyObject_Call
      0
          0.0%
      0
          0.0%
                           35 77.8% PyRun_StringFlags
                0.0%
                            35 77.8% __Pyx_PyObject_Call.constprop.73
      \cap
          0.0%
```

For more information on %crun, see sage.misc.gperftools.

C-level line-by-line profiling: perf (Linux only)

If your code is written in C or in Cython, you can find out line-by-line which are the most costly using perf (included in the Ubuntu package linux-tools).

The easiest way to use it is to run some (very long) computation in Sage, and to type in a console:

```
[user@localhost ~] sudo perf top
```

Select the entry that interests you, and press Enter. The annotate command will show you:

- the CPU instructions
- · the source code
- · the associated time

Note:

- press s to toggle source code view
- press H to cycle through hottest instructions
- press h for help

Alternatively, or if you have no sudo privileges, you can record the statistics of a specific process into a file perf.data from its PID. Then, visualize the result using perf report:

```
[user@localhost ~] perf record -p PID
[user@localhost ~] perf report --vmlinux vmlinux
```

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