

# Structure and Interpretation of Classical Mechanics with Python and Sagemath

Nicky van Foreest

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## PRELIMINARIES

---

### 0.1 README

This is a translation to Python and Sagemath of (most of) the Scheme code of the book ‘Structure and interpretation of classical mechanics’ by Sussman and Wisdom. When referring to *the book*, I mean their book. I expect the reader to read the related parts of the book, and use the Python code to understand the Scheme code of the book (and vice versa). I therefore don’t explain much of the logic of the code in this document. I’ll try to stick to the naming of functions and variables as used in the book. I also try to keep the functional programming approach of the book; consequently, I don’t strive to the most pythonic code possible. To keep the code clean, I never protect functions against stupid input; realize that this is research project, the aim is not to produce a fool-proof software product.

- The file `sicm_sagemath.pdf` shows all code samples together with the output when running the code.
- The directory `org` contains the org files.
- The directory `sage` contains all sage files obtained from tangling the org files.

In the pdf file I tend to place explanations, comments, and observations about the code and the results *above* the code blocks.

I wrote this document in Emacs and Org mode. When developing, I first made a sage file with all code for a specific section of the book. Once everything worked, I copied the code to an Org file and make code blocks. Then I tangled, for instance, generally useful code of `section1.4.org` to `utils1.4.sage` and code specific for Section 1.4 to `section1.4.sage`. A util file does not contain code that will be executed when loading the util file. This way I can load the utils files at later stages.

I found it convenient to test things in a `tests.sage` file. I can edit `tests.sage` within Emacs and see the consequences directly in the sage session by opening a sage session on the command prompt and attaching the session to the file like so:

```
sage: attach("tests.sage")
```

Finally, here are some resources that were helpful to me:

- An online version of the book: <https://tgvaughan.github.io/sicm/>
- An org file of the book with Scheme: <https://github.com/mentat-collective/sicm-book/blob/main/org/chapter001.org>

- A port to Clojure: <https://github.com/sicmutils/sicmutils>
- The Sagemath reference guide: <https://doc.sagemath.org/html/en/reference/>
- Handy tuples: <https://github.com/jtauber/functional-differential-geometry>
- ChatGPT proved to be a great help in the process of becoming familiar with Scheme and Sagemath.
- Some solutions to problems: <https://github.com/hnarayanan/sicm>

In the next sections we provide Python and Sagemath code for background functions that are used, but not defined, in the book.

## 0.2 OUTPUT TO LATEX

We need some tricks to adapt the LATEX output of Sagemath to our liking.

We use `re` to modify LateX strings. I discovered the two `latex` options from this site: [Sage, LateX and Friends](#).

---

```

1 import re
2
3 latex.matrix_delimiters(left='[', right=']')
4 latex.matrix_column_alignment("c")

```

---

Note in passing that the title of the code block shows the file to which the code is tangled, and if a code block is not tangled, the title says "don't tangle".

To keep the formulas short in LATEX, I remove all strings like  $(t)$ , and replace  $\partial x / \partial t$  by  $\dot{x}$ . This is the job of the regular expressions below.

---

```

5 def simplify_latex(s):
6     s = re.sub(r"\\"frac{\\"partial}{\\"partial t}", r"\\"dot ", s)
7     s = re.sub(r"\\"left\((t\\"right)", r"\", s)
8     s = re.sub(
9         r"\\"frac{\\"partial^{\\"{2}}}{\\"partial t\\"^{2}}",
10        r"\\"ddot ",
11        s,
12    )
13    return s

```

---

The function `show_expression` prints expressions to LATEX. There is a caveat, though. When `show_expression` would return a string, org mode (or perhaps Python) adds many escape symbols for the `\` character, which turns out to ruin the LATEX output in an org file. For this reason, I just call `print`; for my purposes (writing these files in emacs and org mode) it works the way I want.

---

```

14 def show_expression(s, simplify=True):
15     s = latex(s)
16     if simplify:
17         s = simplify_latex(s)
18     res = r"\begin{dmath*}"
19     res += "\n" + s + "\n"
20     res += r"\end{dmath*}"
21     print(res)

```

---

### 0.2.1 Printing with org mode

There is a subtlety with respect to printing in org mode and in tangled files. When working in sage files, and running them from the prompt, I call `show(expr)` to have some expression printed to the screen. So, when running Sage from the prompt, I do *not* want to see L<sup>A</sup>T<sub>E</sub>X output. However, when executing a code block in org mode, I *do* want to get L<sup>A</sup>T<sub>E</sub>X output. For this, I could use the book's `show_expression` in the code blocks in the org file. So far so good, but now comes the subtlety. When I *tangle* the code from the org file to a sage file, I don't want to see `show_expression`, but just `show`. Thus, I should use `show` throughout, but in the org mode file, `show` should call `show_expression`. To achieve this, I include the following `show` function in org mode, but I don't *tangle* it to the related sage files.

---

```

22 def show(s, simplify=True):
23     return show_expression(s, simplify)

```

---

## 0.3 THE TUPLE CLASS

The book uses up tuples quite a bit. This code is a copy of `tuples.py` from <https://github.com/jtauber/functional-differential-geometry>. See `tuples.rst` in that repo for further explanations.

---

```

24 """
25 This is a copy of tuples.py from
26 https://github.com/jtauber/functional-differential-geometry.
27 """
28
29 from sage.structure.element import Matrix, Vector
30
31 class Tuple:
32     def __init__(self, *components):
33         self._components = components
34
35     def __getitem__(self, index):

```

```

36     return self._components[index]
37
38     def __len__(self):
39         return len(self._components)
40
41     def __eq__(self, other):
42         if (
43             isinstance(other, self.__class__)
44             and self._components == other._components
45         ):
46             return True
47         else:
48             return False
49
50     def __ne__(self, other):
51         return not (self.__eq__(other))
52
53     def __add__(self, other):
54         if isinstance(self, Tuple):
55             if not isinstance(other, self.__class__) or len(self) != len(
56                 other
57             ):
58                 raise TypeError("can't add incompatible Tuples")
59             else:
60                 return self.__class__(
61                     *(
62                         s + o
63                         for (s, o) in zip(self._components, other._components)
64                     )
65                 )
66         else:
67             return self + other
68
69     def __iadd__(self, other):
70         return self + other
71
72     def __neg__(self):
73         return self.__class__(*(-s for s in self._components))
74
75     def __sub__(self, other):
76         return self + (-other)
77
78     def __isub__(self, other):
79         return self - other
80
81     def __call__(self, **kwargs):
82         return self.__class__(
83             *(
84                 (c(**kwargs) if isinstance(c, Expr) else c)
85                 for c in self._components
86             )
87         )

```

```

88
89     def subs(self, args):
90         # substitute variables with args
91         return self.__class__(*[c.subs(args) for c in self._components])
92
93     def list(self):
94         "convert tuple and its components to one list."
95         result = []
96         for comp in self._components:
97             if isinstance(comp, (Tuple, Matrix, Vector)):
98                 result.extend(comp.list())
99             else:
100                 result.append(comp)
101
102     return result
103
104     def derivative(self, var):
105         "Compute the derivative of all components and put the result in a tuple."
106         return self.__class__
107             *[derivative(comp, var) for comp in self._components]
108

```

---

We have up tuples and down tuples. They differ in the way they are printed.

```

----- ./sage/tuples.sage -----
108 class UpTuple(Tuple):
109     def __repr__(self):
110         return "up({})".format(", ".join(str(c) for c in self._components))
111
112     def _latex_(self):
113         "Print up tuples vertically."
114         res = r"\begin{array}{c}"
115         for comp in self._components:
116             res += r"\begin{array}{c}"
117             res += latex(comp)
118             res += r"\end{array}"
119             res += r"\\"
120         res += r"\end{array}"
121
122     return res
123
124 class DownTuple(Tuple):
125     def __repr__(self):
126         return "down({})".format(", ".join(str(c) for c in self._components))
127
128     def _latex_(self):
129         "Print down tuples horizontally."
130         res = r"\begin{array}{c}"
131         for comp in self._components:
132             res += r"\begin{array}{c}"
133             res += latex(comp)
134             res += r"\end{array}"
135             res += r" & "
136         res += r"\end{array}"

```

```

136     return res
137
138 up = UpTuple
139 down = DownTuple
140
141 up._dual = down
142 down._dual = up

```

---

Here is some functionality to unpack tuples. I don't use it for the moment, but it is provided by the `tuples.py` package that I downloaded from the said github repo.

```

----- ./sage/tuples.sage -----
143 def ref(tup, *indices):
144     if indices:
145         return ref(tup[indices[0]], *indices[1:])
146     else:
147         return tup
148
149
150 def component(*indices):
151     def _(tup):
152         return ref(tup, *indices)
153
154     return _

```

---

## 0.4 FUNCTIONAL PROGRAMMING WITH PYTHON FUNCTIONS

In this section we set up some generic functionality to support the summation, product, and composition of functions:

$$\begin{aligned}(f + g)(x) &= f(x) + g(x), \\ (fg)(x) &= f(x)g(x), \\ (f \circ g)(x) &= f(g(x)).\end{aligned}$$

This is easy to code with recursion.

### 0.4.1 Standard imports

```

----- ./sage/functions.sage -----
155 load("tuples.sage")

```

---

We need to load `functions.sage` to run the examples in the test file.

```

----- ./sage/functions_tests.sage -----
156 load("functions.sage")

```

---

We load `show_expression` to control the L<sup>A</sup>T<sub>E</sub>X output in this org file.

---

don't tangle

---

```
157 load("show_expression.sage")
```

---

### 0.4.2 The Function class

The Function class provides the functionality we need for functional programming.

---

..../sage/functions.sage

---

```
158 class Function:
159     def __init__(self, func):
160         self._func = func
161
162     def __call__(self, *args):
163         return self._func(*args)
164
165     def __add__(self, other):
166         return Function(lambda *args: self(*args) + other(*args))
167
168     def __neg__(self):
169         return Function(lambda *args: -self(*args))
170
171     def __sub__(self, other):
172         return self + (-other)
173
174     def __mul__(self, other):
175         if isinstance(other, Function):
176             return Function(lambda *args: self(*args) * other(*args))
177         return Function(lambda *args: other * self(*args))
178
179     def __rmul__(self, other):
180         return self * other
181
182     def __pow__(self, exponent):
183         if exponent == 0:
184             return Function(lambda x: 1)
185         else:
186             return self * (self ** (exponent - 1))
```

---

The next function decorates a function `f` that returns another function `inner_f`, so that `inner_f` becomes a `Function`.

---

..../sage/functions.sage

---

```
187 def Func(f):
188     def wrapper(*args, **kwargs):
189         return Function(f(*args, **kwargs))
190
191     return wrapper
```

---

Below I include an example to see how to use, and understand, this decorator. Composition is just a recursive call of functions.

---

```

192 @Func
193 def compose(*funcs):
194     if len(funcs) == 1:
195         return lambda x: funcs[0](x)
196     return lambda x: funcs[0](compose(*funcs[1:])(x))

```

---

### 0.4.3 Some standard functions

To use python functions as Functions, use `lambda` like this.

---

```

197 def f(x):
198     return 5 * x
199
200 F = Function(lambda x: f(x))

```

---

The identity is just interesting. Perhaps we'll use it later.

---

```

202 identity = Function(lambda x: x)

```

---

To be able to code things like `(sin + cos)(x)` we need to postpone the application of `sin` and `cos` to their arguments. Therefore we override their definitions.

---

```

203 sin = Function(lambda x: sage.functions.trig.sin(x))
204 cos = Function(lambda x: sage.functions.trig.cos(x))

```

---

We will use quadratic functions often.

---

```

205 from functools import singledispatch
206
207
208 @singledispatch
209 def _square(x):
210     raise TypeError(f"Unsupported type: {type(x)}")
211
212 @_square.register(int)
213 @_square.register(float)
214 @_square.register(Expression)
215 @_square.register(Integer)
216
217 def _(x):
218     return x ^ 2
219
220

```

```

221 @_square.register(Vector)
222 @_square.register(list)
223 @_square.register(tuple)
224 def _(x):
225     v = vector(x)
226     return v.dot_product(v)
227
228
229 @_square.register(Matrix)
230 def _(x):
231     if x.ncols() == 1:
232         return (x.T * x)[0, 0]
233     elif x nrows() == 1:
234         return (x * x.T)[0, 0]
235     else:
236         raise TypeError(
237             f"Matrix must be a row or column vector, got shape {x nrows()}x{x ncols()}"
238         )
239
240
241 square = Function(lambda x: _square(x))

```

---

To use Sagemath functions we make an abbreviation.

```

..... ./sage/functions.sage
242 function = sage.symbolic.function_factory.function

```

---

Now we can make symbolic functions like so.

```

..... ./sage/functions_tests.sage
243 V = Function(lambda x: function("V")(x))

```

---

#### 0.4.4 Examples

```

..... ./sage/functions_tests.sage
244 x, y = var("x y", domain = RR)
245 show((square)(x + y).expand())

```

---

$$x^2 + 2xy + y^2$$

```

..... ./sage/functions_tests.sage
247 show((square + square)(x + y))

```

---

$$2(x + y)^2$$

```

..... ./sage/functions_tests.sage
248 show((square * square)(x))

```

---

$$x^4$$

---

```
249      show((sin + cos)(x))
```

---

$$\cos(x) + \sin(x)$$

---

```
250      show((square + V)(x))
```

---

$$x^2 + V(x)$$

---

```
251      hh = compose(square, sin)
252      show((hh + hh)(x))
```

---

$$2 \sin(x)^2$$

We know that  $2 \sin x \cos x = \sin(2x)$ .

---

```
253      show((2 * (sin * cos)(x) - sin(2 * x)).simplify_full())
```

---

$$0$$

Next, we test differentiation and integration.

---

```
254      show(diff(-compose(square, cos)(x), x))
255      show(integrate((2 * sin * cos)(x), x))
```

---

$$2 \cos(x) \sin(x)$$

$$- \cos(x)^2$$

Arithmetic with symbolic functions works too.

---

```
256      U = Function(lambda x: function("U")(x))
257      V = Function(lambda x: function("V")(x))
```

---



---

```
258      show((U + V)(x))
259      show((V + V)(x))
260      show((V(U(x))))))
261      show((compose(V, U)(x))))
```

---

$$U(x) + V(x)$$

$$2V(x)$$

$$V(U(x))$$

$$V(U(x))$$

---

```
262     def f(x):
263         def g(y):
264             return x * y ^ 2
265
266         return g


---


267 show(f(3)(5))
```

---

75

However, we cannot apply algebraic operations on `f`. For instance, this does not work; it gives `TypeError: unsupported operand type(s) for +: 'function' and 'function'`.

---

```
268 show((f(3) + f(2))(4))
```

---

By decoration with `@Func` we get what we need.

---

```
269 @Func
270 def f(x):
271     def g(y):
272         return x * y ^ 2
273
274     return g


---


275 show((f(3) + f(2))(4))
```

---

80

Indeed:  $(3 + 2) * 4^2 = 80$ .

Decorating with `@Func` is the same as this.

---

```
276 def f(x):
277     def g(y):
278         return x * y ^ 2
279
280     return Function(lambda y: g(y))


---


281 show((f(3) + f(2))(4))
```

---

80

## 0.5 DIFFERENTIATION

### 0.5.1 Standard imports

```

282 _____ ..../sage/differentiation.sage _____
283 load(
284     "functions.sage",
285     "tuples.sage",
286 )
287 _____ ..../sage/differentiation_tests.sage _____
288 load("differentiation.sage")
289 var("t", domain="real")
290 _____ don't tangle _____
291 load("show_expression.sage")
292 
```

### 0.5.2 Examples with matrices, functions and tuples

```

290 _____ ..../sage/differentiation_tests.sage _____
291 _ = var("a b c x y", domain=RR)
292 M = matrix([[a, b], [b, c]])
293 b = vector([a, b])
294 v = vector([x, y])
295 F = 1 / 2 * v * M * v + b * v + c
296 _____ ..../sage/differentiation_tests.sage _____
297 show(F)
298 
```

$$\frac{1}{2}(ax + by)x + ax + \frac{1}{2}(bx + cy)y + by + c$$

```

296 _____ ..../sage/differentiation_tests.sage _____
297 show(F.expand())
298 
```

$$\frac{1}{2}ax^2 + bxy + \frac{1}{2}cy^2 + ax + by + c$$

```

297 _____ ..../sage/differentiation_tests.sage _____
298 show(diff(F, x))
299 
```

$$ax + by + a$$

Repeated differentiation works nicely.

---

```
298 ..../sage/differentiation_tests.sage
show(diff(F, [x, y]))
```

---

$$b$$

This is the Jacobian.

---

```
299 ..../sage/differentiation_tests.sage
show(jacobian(F, [x, y]))
```

---

$$\begin{bmatrix} ax + by + a & bx + cy + b \end{bmatrix}$$

---

```
300 ..../sage/differentiation_tests.sage
show(jacobian(F, v.list())) # convert the column matrix to a list
```

---

$$\begin{bmatrix} ax + by + a & bx + cy + b \end{bmatrix}$$

This expression gives an error.

---

```
301 ..../sage/differentiation_tests.sage
diff(F, v) # v is not a list, but a vector
```

---

To differentiate a Python function we need to provide the arguments to the function.

---

```
302 ..../sage/differentiation_tests.sage
def F(v):
    return 1 / 2 * v * M * v + b * v + c
```

---


---

```
304 ..../sage/differentiation_tests.sage
show(diff(F(v), x)) # add the arguments to F
show(jacobian(F(v), v.list()))
```

---

$$ax + by + a$$

$$\begin{bmatrix} ax + by + a & bx + cy + b \end{bmatrix}$$

The next two examples do not work.

---

```
306 ..../sage/differentiation_tests.sage
don't tangle
jacobian(F, v) # F has no arguments
jacobian(F(v), v) # v is not a list
```

---

The Tuple class supports differentiation.

---

```
308 ..../sage/differentiation_tests.sage
T = up(t, t ^ 2, t ^ 3, sin(3 * t))
309 show(diff(T, t))
```

---

$$\begin{aligned} 1 \\ 2t \\ 3t^2 \\ 3 \cos(3t) \end{aligned}$$

### 0.5.3 Differentiation with respect to time

The function `D` takes a function (of time) as argument, and returns the derivative with respect to time:

$$D(f(\cdot)): t \rightarrow f'(t).$$

---

```
310 @Func
311 def D(f):
312     return lambda t: diff(f(t), t)
313 #return derivative(expr, t)
```

---

Here is an example.

---

```
314 q = Function(lambda t: function("q")(t)) don't tangle
315
316 show(D(q)(t))
```

---

$$\dot{q}$$

### 0.5.4 Differentiation with respect to function arguments

The Euler-Lagrange equations depend on the partial derivative of a Lagrangian  $L$  with respect to  $q$  and  $v$ , and a total derivative with respect to time. Now  $q$  and  $v$  will often be functions of time, so we need to find a way to differentiate with respect to *functions*, like  $q(\cdot)$ , rather than just symbols, like  $x$ . To implement this in Sagemath turned out to be far from easy, at least for me.

First, observe that the Jacobian in Sagemath takes as arguments a function and the variables with respect to which to take the derivatives. So, I tried this first:

---

```
317 q = Function(lambda t: function("q")(t))
```

---

But the next code gives errors saying that the argument  $q$  should be a symbolic function, which it is not.

---

```
318 F = 5 * q + 3 * t don't tangle
319
320 show(diff(F, r)) # does not work
321 show(jacobian(F, [q, t])) # does not work
```

---

To get around this problem, I use the following strategy to differentiate a function  $F$  with respect to functions.

1. Make a list of dummy symbols, one for *each argument* of  $F$  that is *not a symbol*. To understand this in detail, observe that arguments like  $t$  or  $x$  are symbols, but such symbols need not be protected. In other words: we don't have to replace a symbol by another symbol, because Sagemath can already differentiate wrt symbols; it's the other 'things' are the things that have to be replaced by a variable. Thus, arguments like  $q(t)$  that are *not* symbols have to be protected by replacing them with dummy symbols.
2. Replace in  $F$  the arguments by their dummy variables. We use the Sagemath `subs` functionality of Sagemath to substitute the dummy variables for the functions. Now there is one further problem: `subs` does not work on lists or tuples. However, `subs` *does work* on vectors and matrices. Therefore, we cast all relevant lists to vectors, which suffices for our goal.
3. Take the Jacobian of  $F$  with respect to the dummy symbols. We achieve this by substituting the dummy symbols in the vector of arguments and the vector of variables.
4. Invert: Replace in the final result the dummy symbols by their related arguments.

We use `id(v)` to create a unique variable name for each dummy variable and store the mapping from the functions to the dummy variables in a dictionary `subs`. (As these are internal names, the actual variable names are irrelevant; as long as they are unique, it's OK.)

We know from the above that `jacobian` expects a *list* with the variables with respect to which to differentiate. Therefore, we turn the vector with substituted variables to a list.

---

```

322 def Jacobian(F):
323     def f(args, vrs):
324         if isinstance(args, (list, tuple)):
325             args = vector(args)
326         if isinstance(vrs, (list, tuple)):
327             vrs = vector(vrs)
328         subs = {
329             v: var(f"v{id(v)}", domain=RR)
330             for v in args.list()
331             if not v.is_symbol()
332         }
333         result = jacobian(F(args.subs(subs)), vrs.subs(subs).list())
334         inverse_subs = {v: k for k, v in subs.items()}
335         return result.subs(inverse_subs)
336
337     return f

```

---

Here are some examples to see how to use this Jacobian. Note that Jacobian expects the arguments and variables to be *lists*, or list like. As a result, in the function `F` we have to unpack the list.

---

```

338 v = var("v", domain=RR)
339
340
341 def F(v):
342     r, t = v.list()
343     return 5 * r ^ 3 + 3 * t ^ 2 * r
344
345
346 show(Jacobian(F)([v, t], [t]))
347 show(Jacobian(F)([v, t], [v, t]))

```

---

$$[ 6tv ]$$

$$[ 3t^2 + 15v^2 \quad 6tv ]$$

This works. Now we try the same with a function like argument. Recall, `v` must be a list for `partial` on which gradient depends.

---

```

348 q = Function(lambda t: function("q")(t))
349 v = [q(t), t]
350 show(Jacobian(F)(v, v))

```

---

$$[ 3t^2 + 15q^2 \quad 6tq ]$$

### 0.5.5 Gradient and Hessian

Next we build the gradient. We can use Sagemath's `jacobian`, but as is clear from above, we need to indicate explicitly the variable names with respect to which to differentiate. Moreover, we like to be able to take the gradient with respect to literal functions. Thus, we use the `Jacobian` defined above.

One idea for the gradient is like this. However, this does not allow to use `gradient` as a function in functional composition.

---

```

351 def gradient(F, v):
352     return Jacobian(F)(v, v).T

```

---

We therefore favor the next implementation. BTW, note that the gradient is a vector in a tangent space, hence it is column vector. For that reason we transpose the Jacobian.

---

```

353 def gradient(F):
354     return lambda v: Jacobian(F)(v, v).T

```

---

---

```
355 ..../sage/differentiation_tests.sage
show(gradient(F)(v))
```

---

$$\begin{bmatrix} 3t^2 + 15q^2 \\ 6tq \end{bmatrix}$$

When differentiating a symbolic function, wrap such a function in a Function.

---

```
356 ..../sage/differentiation_tests.sage
U = Function(lambda x: function("U")(square(x)))
357 show(gradient(U)(v))
```

---

$$\begin{bmatrix} 2qD_0(U)(t^2 + q^2) \\ 2tD_0(U)(t^2 + q^2) \end{bmatrix}$$

The Hessian can now be defined as the composition of the gradient with itself.

---

```
358 ..../sage/differentiation.sage
def Hessian(F):
359     return lambda v: compose(gradient, gradient)(F)(v)
360 ..../sage/differentiation_tests.sage
show(Hessian(F)(v))
```

---

$$\begin{bmatrix} 30q & 6t \\ 6t & 6q \end{bmatrix}$$

### 0.5.6 Differentiation with respect to slots

To follow the notation of the book, we need to define a python function that computes partial derivatives with respect to the slot of a function; for example, in  $\partial_1 L$  the 1 indicates that the partial derivatives are supposed to be taken wrt the coordinate variables. The Jacobian function built above allows us a very simple solution. Note that we return a Function so that we can use this operator in functional composition if we like.

---

```
361 ..../sage/differentiation.sage
362 @Func
363 def partial(f, slot):
364     def wrapper(local):
365         if slot == 0:
366             selection = [time(local)]
367         elif slot == 1:
368             selection = coordinate(local)
369         elif slot == 2:
370             selection = velocity(local)
371     return Jacobian(f)(local, selection)
372
373 return wrapper
```

---

The main text contains many examples.

## CHAPTER 1

---

### 1.4 COMPUTING ACTIONS

#### 1.4.1 Standard setup

I create an Org file for each separate section of the book; for this section it's `section1.4.org`. Code that is useful for later sections is tangled to `utils1.4.sage` and otherwise to `section1.4.sage`. This allows me to run the sage scripts on the prompt. Note that the titles of the code blocks correspond to the file to which the code is written when tangled.

---

```
373 import numpy as np
374
375 load("functions.sage", "differentiation.sage", "tuples.sage")
```

---

BTW, don't do `from sage.all import *` because that will lead to name space conflicts, for instance with the Gamma function which we define below.

---

```
376 load("utils1.4.sage")
377
378 t = var("t", domain="real")
```

---

The next module is used for nice printing in org mode files; it should only be loaded in org mode files.

---

```
379 load("show_expression.sage")
```

---

#### 1.4.2 The Lagrangian for a free particle.

The function `L_free_particle` takes `mass` as an argument and returns the (curried) function `Lagrangian` that takes a `local` tuple as an argument.

---

```
380 def L_free_particle(mass):
381     def Lagrangian(local):
382         v = velocity(local)
383         return 1 / 2 * mass * square(v)
384
385     return Lagrangian
```

---

For the next step, we need a *literal functions* and *coordinate paths*.

### 1.4.3 Literal functions

A `literal_function` maps the time  $t$  to a coordinate or velocity component of the path, for instance,  $t \rightarrow x(t)$ . Since we need to perform arithmetic with literal functions, see below for some examples, we encapsulate it in a `Function`.

---

```
386     _____ . /sage/utils1.4.sage _____
387 @Func
388 def literal_function(name):
389     return lambda t: function(name)(t)
```

---

It's a function.

---

```
390 _____ don't tangle _____
391 x = literal_function("x")
392 print(x)
```

---

`<__main__.Function object at 0x71122066e470>`

Here are some operations on `x`.

---

```
391 _____ don't tangle _____
392 show(x(t))
393 show((x+x)(t))
394 show(square(x)(t))
```

---

Note that, to keep the notation brief, the  $t$  is suppressed in the L<sup>A</sup>T<sub>E</sub>X output.

### 1.4.4 Paths

We will represent coordinate path functions  $q$  and velocity path functions  $v$  as functions that map time to vectors. Thus, `column_path` returns a function of time, not yet a path. We also need to perform arithmetic on paths, like  $3q$ , therefore we encapsulate the path in a `Function`.

---

```
394     _____ . /sage/utils1.4.sage _____
395 @Func
396 def column_path(lst):
397     #return lambda t: vector([l(t) for l in lst])
398     return lambda t: column_matrix([l(t) for l in lst])
```

---

---

```

398 q = column_path(          don't tangle
399   [
400     literal_function("x"),
401     literal_function("y"),
402   ]
403 )

```

---

Here is an example to see how to use `q`.

---

```

404 show(q(t))          don't tangle

```

---

$$\begin{bmatrix} x \\ y \end{bmatrix}$$


---

---

```

405 show((q + q)(t))    don't tangle

```

---

$$\begin{bmatrix} 2x \\ 2y \end{bmatrix}$$


---

---

```

406 show((2 * q)(t))    don't tangle

```

---

$$\begin{bmatrix} 2x \\ 2y \end{bmatrix}$$


---

---

```

407 show((q * q)(t))    don't tangle

```

---

#### 1.4.5 Gamma function

The Gamma function lifts a coordinate path to a function that maps time to a local tuple of the form  $(t, q(t), v(t), \dots)$ . That is,

$$\begin{aligned}\Gamma[q](\cdot) &= (\cdot, q(\cdot), v(\cdot), \dots), \\ \Gamma[q](t) &= (t, q(t), v(t), \dots).\end{aligned}$$

To follow the conventions of the book, we use an up tuple for `Gamma`. However, I don't build the coordinate path nor the velocity as up tuples because I find Sagemath vectors more convenient.

$\Gamma$  just receives  $q$  as an argument. Then it computes the velocity  $v = Dq$ , from which the acceleration follows recursively as  $a = Dv, \dots$ . Recall that `D` computes the derivative (wrt time) of a function that depends on time.

When  $n = 3$ , it returns a function of time that produces the first three elements of the local tuple  $(t, q(t), v(t))$ . This is the default. Once all derivatives are computed, we convert the result to a function that maps time to an up tuple.

---

```
..... ./sage/utils1.4.sage
408 def Gamma(q, n=3):
409     if n < 2:
410         raise ValueError("n must be > 1")
411     Dq = [q]
412     for k in range(2, n):
413         Dq.append(Dq[-1])
414     return lambda t: up(t, *[v(t) for v in Dq])
```

---

When applying `Gamma` to a path, we get this.

---

```
..... don't tangle
415 local = Gamma(q)(t)
416 show(local)
```

---

$$\begin{bmatrix} t \\ x \\ y \\ \dot{x} \\ \dot{y} \end{bmatrix}$$

We can include the acceleration too.

---

```
..... don't tangle
417 show(Gamma(q, 4)(t))
```

---

$$\begin{bmatrix} t \\ x \\ y \\ \dot{x} \\ \ddot{x} \\ \dot{y} \\ \ddot{y} \end{bmatrix}$$

Finally, here are some projections operators from the local tuple to superspaces.

---

```
..... ./sage/utils1.4.sage
418 time = Function(lambda local: local[0])
419 coordinate = Function(lambda local: local[1])
420 velocity = Function(lambda local: local[2])
..... don't tangle
421 show(compose(velocity, Gamma(q))(t))
```

---

$$\begin{bmatrix} \dot{x} \\ \dot{y} \end{bmatrix}$$

### 1.4.6 Continuation with the free particle.

Now we know how to build literal functions and  $\Gamma$ , we can continue with the Lagrangian of the free particle.

```
422 q = column_path(           .../sage/section1.4.sage
423   [
424     literal_function("x"),
425     literal_function("y"),
426     literal_function("z"),
427   ]
428 )
```

---

```
429 show(q(t))           .../sage/section1.4.sage
```

---


$$\begin{bmatrix} x \\ y \\ z \end{bmatrix}$$


---

```
430 show(D(q)(t))       .../sage/section1.4.sage
```

---


$$\begin{bmatrix} \dot{x} \\ \dot{y} \\ \dot{z} \end{bmatrix}$$


---

```
431 show(Gamma(q)(t))  .../sage/section1.4.sage
```

---


$$\begin{bmatrix} t \\ x \\ y \\ z \\ \dot{x} \\ \dot{y} \\ \dot{z} \end{bmatrix}$$

The Lagrangian of a free particle with mass  $m$  applied to the path  $\text{Gamma}$  gives this. Our first implementation is like this:  $L(\Gamma[q](t))$ , that is,  $\Gamma[q](t)$  makes a local tuple, and this is given as argument to  $L$ .

```
432 load("functions.sage")    .../sage/section1.4.sage
433 m = var('m', domain='positive')
434 show(L_free_particle(m)(Gamma(q)(t)))
```

---

$$\frac{1}{2} (\dot{x}^2 + \dot{y}^2 + \dot{z}^2) m$$

Here is the implementation of the book:  $(L \circ \Gamma[q])(t)$ , that is,  $L \circ \Gamma[q]$  is a function that depends on  $t$ . Note how the brackets are placed after `Gamma(q)`.

---

```
435     show(compose(L_free_particle(m), Gamma(q))(t))
```

---

$$\frac{1}{2} (\dot{x}^2 + \dot{y}^2 + \dot{z}^2) m$$

We now compute the integral of Lagrangian  $L$  along the path  $q$ , but for this we need a function to carry out 1D integration (along time in our case). Of course, Sagemath already supports a definite integral in a library.

---

```
436     from sage.symbolic.integration import definite_integral
```

---

I don't like to read  $dt$  at the end of the integral because  $dt$  reads like the product of the variables  $d$  and  $t$ . Instead, I prefer to read  $dt$ ; for this reason I overwrite the LATEX formatting of `definite_integral`.

---

```
437 def integral_latex_format(*args):
438     expr, var, a, b = args
439     return (
440         fr"\int_{{{{a}}}}^{{{{b}}}} "
441         + latex(expr)
442         + r"\, \text{d}\mathbf{{{d}}}\, "
443         + latex(var)
444     )
445
446
447 definite_integral._print_latex_ = integral_latex_format
```

---

Here is the action along a generic path  $q$ .

---

```
448 T = var("T", domain="positive")
449
450 def Lagrangian_action(L, q, t1, t2):
451     return definite_integral(compose(L, Gamma(q))(t), t, t1, t2)
452
453 show(Lagrangian_action(L_free_particle(m), q, 0, T))
```

---

$$\frac{1}{2} m \left( \int_0^T \dot{x}^2 dt + \int_0^T \dot{y}^2 dt + \int_0^T \dot{z}^2 dt \right)$$

To get a numerical answer, we take the test path of the book. Below we'll do some arithmetic with `test_path`; therefore we encapsulate it in a Function.

---

```
454 test_path = Function(lambda t: vector([4 * t + 7, 3 * t + 5, 2 * t + 1]))
455 show(Lagrangian_action(L_free_particle(mass=3), test_path, 0, 10))
```

---

435

Let's try a harder path. We don't need this later, so the encapsulation in `Function` is not necessary.

---

```
456 hard_path = lambda t: vector([4 * t + 7, 3 * t + 5, 2 * exp(-t) + 1])
457
458 result = Lagrangian_action(L_free_particle(mass=3), hard_path, 0, 10)
459 show(result)
460 show(float(result))
```

---

$$3(125e^{20} - 1)e^{(-20)} + 3$$

377.9999999938165

The value of the integral is different from 435 because the end points of this harder path are not the same as the end points of the test path.

#### 1.4.7 Path of minimum action

First some experiments to see whether my code works as intended.

---

```
461 @Func
462 def make_eta(nu, t1, t2):
463     return lambda t: (t - t1) * (t - t2) * nu(t)
464
465
466 nu = Function(lambda t: vector([sin(t), cos(t), t ^ 2]))
467
468 show((1 / 3 * make_eta(nu, 3, 4) + test_path)(t))
```

---

$$\left(\frac{1}{3}(t-3)(t-4)\sin+4t+7, \frac{1}{3}(t-3)(t-4)\cos+3t+5, \frac{1}{3}(t-3)(t-4)t^2+2t+1\right)$$

In the next code, I add the `n()` to force the result to a floating point number. (Without this, the result is a long expression with lots of cosines and sines.)

---

```
469 def varied_free_particle_action(mass, q, nu, t1, t2):
470     eta = make_eta(nu, t1, t2)
471
```

---

```

472     def f(eps):
473         return Lagrangian_action(L_free_particle(mass), q + eps * eta, t1, t2).n()
474
475     return f
476
477 show(varied_free_particle_action(3.0, test_path, nu, 0.0, 10.0)(0.001))

```

---

436.291214285714

By comparing our result with that of the book, we see we are still on track.  
Now use Sagemath's `find_local_minimum` to minimize over  $\epsilon$ .

```

.../sage/section1.4.sage
478 res = find_local_minimum(
479     varied_free_particle_action(3.0, test_path, nu, 0.0, 10.0), -2.0, 1.0
480 )
481 show(res)

```

---

(435.000000000000,0.0)

We see that the optimal value for  $\epsilon$  is 0, and we retrieve our earlier value of the Lagrangian action.

#### 1.4.8 Finding minimal trajectories

The `make_path` function uses a Lagrangian polynomial to interpolate a given set of data.

```

.../sage/utils1.4.sage
482 def Lagrangian_polynomial(ts, qs):
483     return RR['x'].lagrange_polynomial(list(zip(ts, qs)))

```

---

While a Lagrangian polynomial gives an excellent fit on the fitted points, its behavior in between these points can be quite wild. Let us test the quality of the fit before using this interpolation method. From the book we know we need to fit  $\cos(t)$  on  $t \in [0, \pi/2]$ , so let us try this first before trying to find the optimal path for the harmonic Lagrangian. Since  $\cos^2 x + \sin^2 x = 1$ , we can use this relation to check the quality of derivative of the fitted polynomial at the same time. The result is better than I expected.

```

.../sage/section1.4.sage
484 ts = np.linspace(0, pi / 2, 5)
485 qs = [cos(t).n() for t in ts]
486 lp = Lagrangian_polynomial(ts, qs)
487 ts = np.linspace(0, pi / 2, 20)
488 Cos = [lp(x=t).n() for t in ts]
489 Sin = [lp.derivative(x)(x=t).n() for t in ts]
490 Zero = [abs(Cos[i] ^ 2 + Sin[i] ^ 2 - 1) for i in range(len(ts))]
491 show(max(Zero))

```

---

In the function `make_path` we use numpy's `linspace` instead of the linear interpolants of the book. Note that the coordinate paths above are column-vector functions, so `make_path` should return the same type.

---

```
492     def make_path(t0, q0, t1, q1, qs):
493         ts = np.linspace(t0, t1, len(qs) + 2)
494         qs = np.r_[q0, qs, q1]
495         return lambda t: vector([Lagrangian_polynomial(ts, qs)(t)])
```

---

Here is the harmonic Lagrangian.

---

```
496     def L_harmonic(m, k):
497         def Lagrangian(local):
498             q = coordinate(local)
499             v = velocity(local)
500             return (1 / 2) * m * square(v) - (1 / 2) * k * square(q)
501
502         return Lagrangian
```

---



---

```
503     def parametric_path_action(Lagrangian, t0, q0, t1, q1):
504         def f(qs):
505             path = make_path(t0, q0, t1, q1, qs=qs)
506             return Lagrangian_action(Lagrangian, path, t0, t1)
507
508         return f
```

---

Let's try this on the path  $\cos(t)$ . The intermediate values `qs` will be optimized below, whereas `q0` and `q1` remain fixed. Thus, we strip the first and last element of `linspace` to make `qs`. The result tells us what we can expect for the minimal value for the integral over the Lagrangian along the optimal path.

---

```
509     t0, t1 = 0, pi / 2
510     q0, q1 = cos(t0), cos(t1)
511     T = np.linspace(0, pi / 2, 5)
512     initial_qs = [cos(t).n() for t in T][1:-1]
513     parametric_path_action(L_harmonic(m=1, k=1), t0, q0, t1, q1)(initial_qs)
```

---

What is the quality of the path obtained by the Lagrangian interpolation? (Recall that a path is a vector; to extract the value of the element that corresponds to the path, we need to write `best_path(t=t)[0]`.)

---

```
514     def find_path(Lagrangian, t0, q0, t1, q1, n):
515         ts = np.linspace(t0, t1, n)
516         initial_qs = np.linspace(q0, q1, n)[1:-1]
```

---

```

517     minimizing_qs = minimize(
518         parametric_path_action(Lagrangian, t0, q0, t1, q1),
519         initial_qs,
520     )
521     return make_path(t0, q0, t1, q1, minimizing_qs)
522
523 best_path = find_path(L_harmonic(m=1, k=1), t0=0, q0=1, t1=pi / 2, q1=0, n=5)
524 result = [
525     abs(best_path(t)[0].n() - cos(t).n()) for t in np.linspace(0, pi / 2, 10)
526 ]
527 show(max(result))

```

0.000172462354236957

Great. All works!

Finally, here is a plot of the Lagrangian as a function of  $q(t)$ .

```

----- ./sage/section1.4.sage -----
528 T = np.linspace(0, pi / 2, 20)
529 q = lambda t: vector([cos(t)])
530 lvalues = [L_harmonic(m=1, k=1)(Gamma(q)(t))(t=ti).n() for ti in T]
531 points = list(zip(ts, lvalues))
532 plot = list_plot(points, color="black", size=30)
533 plot.axes_labels(["$t$", "$L$"])
534 plot.save("../figures/Lagrangian.png", figsize=(4, 2))

```

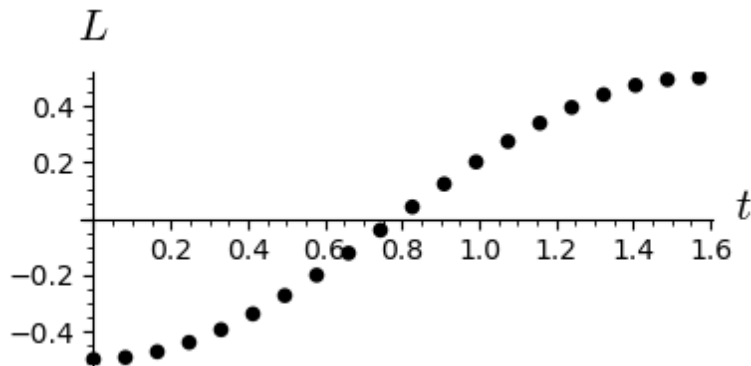


Figure 1.1: The harmonic Lagrangian as a function of the optimal path  $q(t) = \cos t$ ,  $t \in [0, \pi/2]$ .

## 1.5 THE EULER-LAGRANGE EQUATIONS

### 1.5.1 Standard imports

```

----- ./sage/utils1.5.sage -----
535 load("utils1.4.sage")

```

---

```

536   _____ ./.sage/section1.5.sage _____
537   load("utils1.5.sage")
538   t = var("t", domain="real")
_____  

539   _____ don't tangle _____
load("show_expression.sage")
_____

```

---

### 1.5.2 Derivation of the Lagrange equations

#### Harmonic oscillator

Here is a test on the harmonic oscillator.

---

```

540   _____ ./.sage/section1.5.sage _____
541   load("utils1.4.sage")
542   k, m = var('k m', domain="positive")
543   q = column_path([literal_function("x")])
_____  

543   _____ ./.sage/section1.5.sage _____
L = L_harmonic(m, k)
show(L(Gamma(q)(t)))
_____

```

---

$$-\frac{1}{2}kx^2 + \frac{1}{2}m\dot{x}^2$$

We can apply  $\partial_1 L$  and  $\partial_2 L$  to a configuration path  $q$  that we lift to a local tuple by means of  $\Gamma$ . Realize therefore that  $\text{partial}(L_{\text{harmonic}}(m, k), 1)$  maps a local tuple to a real number, and  $\text{Gamma}(q)$  maps a time  $t$  to a local tuple. The next code implements  $\partial_1 L(\Gamma(q)(t))$  and  $\partial_2 L(\Gamma(q)(t))$ . (Check how the brackets are organized.)

---

```

545   _____ ./.sage/section1.5.sage _____
show(partial(L, 1)(Gamma(q)(t)))
_____

```

---

$$[ -kx ]$$

---

```

546   _____ ./.sage/section1.5.sage _____
show(partial(L, 2)(Gamma(q)(t)))
_____

```

---

$$[ m\dot{x} ]$$

Here are the same results, but now with functional composition.

$$(\partial_1 L \circ \Gamma(q))(t), \quad (\partial_2 L \circ \Gamma(q))(t).$$

---

```
547 _____ ..../sage/section1.5.sage _____
548 show(compose(partial(L, 1), Gamma(q))(t))
548 show(compose(partial(L, 2), Gamma(q))(t))
```

---

$$[ -kx ]$$

$$[ m\dot{x} ]$$

These results are functions of  $t$ , so we can take the derivative with respect to  $t$ , which forms the last step to check before building the Euler-Lagrange equations. To understand this, note the following function mappings, where we write  $t$  for time,  $l$  for a local tuple,  $v$  a velocity-like vector, and  $a$  an acceleration-like vector:

$$\begin{aligned}\Gamma[q] &: t \rightarrow l, \\ \partial_2 L &: l \rightarrow v \\ \partial_2 L \circ \Gamma[q] &: t \rightarrow v \\ D(v) &: t \rightarrow a \\ D(\partial_2 L \circ \Gamma[q]) &: t \rightarrow a.\end{aligned}$$

In more classical notation, we compute this:

$$\frac{d}{dt} \left( \frac{\partial}{\partial \dot{q}} L(\Gamma(q)) \right)(t)$$

---

```
549 _____ ..../sage/section1.5.sage _____
549 show(D(compose(partial(L, 2), Gamma(q)))(t))
```

---

$$[ m\ddot{x} ]$$

There we are! We can now try the other examples of the book.

### *Orbital motion*

---

```
550 _____ ..../sage/section1.5.sage _____
550 q = column_path([literal_function("xi"), literal_function("eta")])
```

---

```
551 _____ ..../sage/section1.5.sage _____
551 var("mu", domain="positive")
552
553 def L_orbital(m, mu):
554     def Lagrangian(local):
555         q = coordinate(local)
556         v = velocity(local)
557         return (1 / 2) * m * square(v) + mu / sqrt(square(q))
558
559     return Lagrangian
```

---

---

```
560 L = L_orbital(m, mu)
561 show(L(Gamma(q)(t)))
```

---


$$\frac{1}{2} (\dot{\eta}^2 + \dot{\xi}^2) m + \frac{\mu}{\sqrt{\eta^2 + \xi^2}}$$


---

```
562 show(partial(L, 1)(Gamma(q)(t)))
```

---


$$\begin{bmatrix} -\frac{\mu\xi}{(\eta^2+\xi^2)^{\frac{3}{2}}} & -\frac{\mu\eta}{(\eta^2+\xi^2)^{\frac{3}{2}}} \end{bmatrix}$$


---

```
563 show(partial(L, 2)(Gamma(q)(t)))
```

---


$$\begin{bmatrix} m\ddot{\xi} & m\ddot{\eta} \end{bmatrix}$$

*An ideal planar pendulum, Exercise 1.9.a of the book*

We need a new path in terms of  $\theta$  and  $\dot{\theta}$ .

---

```
564 q = column_path([literal_function("theta")])
```

---

Here is the Lagrangian. Recall that the coordinates of the space form a vector. Here, `theta` is the only element of the vector, which we can extract by considering element 0. For `thetadot` we don't have to do this since we consider  $\dot{\theta}^2$ , and the `square` function accepts vectors as input and returns a real. However, for reasons of consistency, we choose to do this nonetheless.

---

```
565 var("m g l", domain="positive")
566
567
568 def L_planar_pendulum(m, g, l):
569     def Lagrangian(local):
570         theta = coordinate(local).list()[0]
571         theta_dot = velocity(local).list()[0]
572         T = (1 / 2) * m * l ^ 2 * square(theta_dot)
573         V = m * g * l * (1 - cos(theta))
574         return T - V
575
576     return Lagrangian
```

---

```
577 L = L_planar_pendulum(m, g, l)
578 show(L(Gamma(q)(t)))
```

---

---

```
579 show(partial(L, 1)(Gamma(q)(t)))
```

---



---

```
580 show(partial(L, 2)(Gamma(q)(t)))
```

---

*Henon Heiles potential, Exercise 1.9.b of the book*

As the potential depends on the  $x$  and  $y$  coordinate separately, we need to unpack the coordinate vector.

---

```
581 def L_Henon_Heiles(m):
582     def Lagrangian(local):
583         x, y = coordinate(local).list()
584         v = velocity(local)
585         T = (1 / 2) * square(v)
586         V = 1 / 2 * (square(x) + square(y)) + square(x) * y - y**3 / 3
587         return T - V
588
589     return Lagrangian
```

---



---

```
590 L = L_Henon_Heiles(m)
591 q = column_path([literal_function("x"), literal_function("y")])
592 show(L(Gamma(q)(t)))
```

---

$$-x^2y + \frac{1}{3}y^3 - \frac{1}{2}x^2 - \frac{1}{2}y^2 + \frac{1}{2}\dot{x}^2 + \frac{1}{2}\dot{y}^2$$

---

```
593 show(partial(L, 1)(Gamma(q)(t)))
```

---

$$\begin{bmatrix} -2xy - x & -x^2 + y^2 - y \end{bmatrix}$$

---

```
594 show(partial(L, 2)(Gamma(q)(t)))
```

---

$$\begin{bmatrix} \dot{x} & \dot{y} \end{bmatrix}$$

*Motion on the 2d sphere, Exercise 1.9.c of the book*

---

```
595 var('R', domain="positive") .../sage/section1.5.sage
596
597
598 def L_sphere(m, R):
599     def Lagrangian(local):
600         theta, phi = coordinate(local).list()
601         alpha, beta = velocity(local).list()
602         L = m * R * (square(alpha) + square(beta * sin(theta))) / 2
603         return L
604
605     return Lagrangian
```

---



---

```
606 q = column_path([literal_function("phi"), literal_function("theta")])
607 L = L_sphere(m, R)
608
609 show(L(Gamma(q)(t)))
```

---

$$\frac{1}{2} (\sin(\phi)^2 \dot{\theta}^2 + \dot{\phi}^2) Rm$$

---

```
610 show(partial(L, 1)(Gamma(q)(t)))
```

---

$$[ Rm \cos(\phi) \sin(\phi) \dot{\theta}^2 \quad 0 ]$$

---

```
611 show(partial(L, 2)(Gamma(q)(t)))
```

---

$$[ Rm\dot{\phi} \quad Rm \sin(\phi)^2 \dot{\theta} ]$$

*Higher order Lagrangians*

I recently read the books of Larry Susskind on the theoretical minimum for physics. He claims that Lagrangians up to first order derivatives suffice to understand nature, so I skip this part.

### 1.5.3 Computing Lagrange's equation

The Euler-Lagrange equations are simple to implement now that we have a good function for computing partial derivatives.

### The Euler Lagrange Equations

We work in steps to see how all components tie together.

---

```

612 q = column_path(
613     [
614         literal_function("x"),
615         literal_function("y"),
616     ]
617 )
618
619 L = L_free_particle(m)
620 show(compose(partial(L, 1), Gamma(q))(t))
621 show(compose(partial(L, 2), Gamma(q))(t))
622 show(D(compose(partial(L, 2), Gamma(q)))(t))
623 show(
624     (D(compose(partial(L, 2), Gamma(q))) - compose(partial(L, 1), Gamma(q)))(t)
625 )

```

---

$$\begin{bmatrix} 0 & 0 \end{bmatrix}$$

$$\begin{bmatrix} m\dot{x} & m\dot{y} \end{bmatrix}$$

$$\begin{bmatrix} m\ddot{x} & m\ddot{y} \end{bmatrix}$$

$$\begin{bmatrix} m\ddot{x} & m\ddot{y} \end{bmatrix}$$

The last step forms the Euler-Lagrange equation, which we can now implement as a function.

---

```

626 def Lagrange_equations(L):
627     def f(q):
628         return D(compose(partial(L, 2), Gamma(q))) - compose(
629             partial(L, 1), Gamma(q)
630         )
631
632     return f

```

---

### The free particle

We compute the Lagrange equation for a path linear in  $t$  for the Lagrangian of a free particle..

---

```

633 var("a b c a0 b0 c0", domain="real")
634 test_path = lambda t: column_matrix([a * t + a0, b * t + b0, c * t + c0])

```

---

Note that if we do not provide the argument  $t$  to  $l\_eq$  we receive a function instead of vector.

---

```
635 l_eq = Lagrange_equations(L_free_particle(m))(test_path)
636 show(l_eq(t))
```

---

$$\begin{bmatrix} 0 & 0 & 0 \end{bmatrix}$$

This is correct since a free particle is not moving in a potential field, hence only depends on the velocity but not the coordinates of the path. But since the velocity is linear in  $t$ , all components along the test path become zero.

Here are the EL equations for a generic 1D path.

---

```
637 q = column_path([literal_function("x")])
638 l_eq = Lagrange_equations(L_free_particle(m))(q)
639 show(l_eq(t))
```

---

$$\begin{bmatrix} m\ddot{x} \end{bmatrix}$$

Equating this to (0) shows that the solution of these differential equations is linear in  $t$ .

### *The harmonic oscillator*

---

```
640 var("A phi omega", domain="real")
641 assume(A > 0)
642 proposed_path = lambda t: vector([A * cos(omega * t + phi)])
```

---

Lagrange\_equations returns a matrix whose elements correspond to the components of the configuration path  $q$ .

---

```
644 l_eq = Lagrange_equations(L_harmonic(m, k))(proposed_path)(t)
645 show(l_eq)
```

---

$$\begin{bmatrix} -Am\omega^2 \cos(\omega t + \phi) + Ak \cos(\omega t + \phi) \end{bmatrix}$$

To obtain the contents of this  $1 \times 1$  matrix, we take the element  $[0][0]$ .

---

```
646 show(l_eq[0][0])
```

---

$$-Am\omega^2 \cos(\omega t + \phi) + Ak \cos(\omega t + \phi)$$

Let's factor out the cosine.

---

```
647 show(l_eq[0, 0].factor())
```

---

$$-(m\omega^2 - k) A \cos(\omega t + \phi)$$

*Kepler's third law*

Recall that to unpack the coordinates, we have to convert the vector to a Python list.

---

```
648 var("G m m1 m2", domain="positive")  
649  
650  
651 def L_central_polar(m, V):  
652     def Lagrangian(local):  
653         r, phi = coordinate(local).list()  
654         rdot, phidot = velocity(local).list()  
655         T = 1 / 2 * m * (square(rdot) + square(r * phidot))  
656         return T - V(r)  
657  
658     return Lagrangian  
659  
660  
661 def gravitational_energy(G, m1, m2):  
662     def f(r):  
663         return -G * m1 * m2 / r  
664  
665     return f
```

---

```
666 q = column_path([literal_function("r"), literal_function("phi")])  
667 V = gravitational_energy(G, m1, m2)  
668 L = L_central_polar(m, V)  
669 show(L(Gamma(q)(t)))
```

---


$$\frac{1}{2} (r^2 \dot{\phi}^2 + \dot{r}^2) m + \frac{Gm_1m_2}{r}$$


---

```
670 l_eq = Lagrange_equations(L)(q)(t)
```

---

```
671 show(l_eq[0, 1] == 0)
```

---

$$mr^2\ddot{\phi} + 2mr\dot{\phi}\dot{r} = 0$$

In this equation, let's divide by  $mr$  to get  $r\ddot{\phi} + 2\dot{\phi}\dot{r} = 0$ , which is equal to  $\partial_t(\dot{\phi}r^2) = 0$ . This implies that  $\dot{\phi}r^2 = C$ , i.e., a constant. If  $r \neq 0$  and constant, which we should assume according to the book, then we see that  $\dot{\phi}$  is constant, so the two bodies rotate with constant angular speed around each other.

What can we say about the other equation?

---

```
672 show(l_eq[0, 0] == 0)
```

---

$$-mr\dot{\phi}^2 + m\ddot{r} + \frac{Gm_1m_2}{r^2} = 0$$

As  $r$  is constant according to the book,  $\ddot{r} = 0$ . By dividing by  $m := m_1m_2/(m_1 + m_2)$ , this equation reduces to  $r^3\dot{\phi}^2 = G(m_1 + m_2)$ , which is the form we were to find according to the exercise.

## 1.6 HOW TO FIND LAGRANGIANS

### 1.6.1 Standard imports

```
673   _____ .. / sage / utils1.6.sage _____
load("utils1.5.sage")  

674   _____ .. / sage / section1.6.sage _____
load("utils1.6.sage")  

675   _____ don't tangle _____
load("show_expression.sage")  

_____
```

### 1.6.2 Constant acceleration

We start with a point in a uniform gravitational field.

```
676   _____ .. / sage / utils1.6.sage _____
var("t", domain="real")
var("g m", domain="positive")
678
679
680 def L_uniform_acceleration(m, g):
681     def wrap_L_unif(local):
682         x, y = coordinate(local).list()
683         v = velocity(local)
684         T = 1 / 2 * m * square(v)
685         V = m * g * y
686         return T - V
687
688     return wrap_L_unif  

689   _____ .. / sage / section1.6.sage _____
q = column_path([literal_function("x"), literal_function("y")])
l_eq = Lagrange_equations(L_uniform_acceleration(m, g))(q)
show(l_eq(t))  

_____
```

$$\begin{bmatrix} m\ddot{x} & gm + m\ddot{y} \end{bmatrix}$$

### 1.6.3 Central force field

---

```
.../sage/utils1.6.sage
692 def L_central_rectangular(m, U):
693     def Lagrangian(local):
694         q = coordinate(local)
695         v = velocity(local)
696         T = 1 / 2 * m * square(v)
697         return T - U(sqrt(square(q)))
698
699     return Lagrangian
```

---

Let us first try this on a concrete potential function.

---

```
.../sage/section1.6.sage
700 def U(r):
701     return 1 / r
...
702 show(Lagrange_equations(L_central_rectangular(m, U))(q)(t))
```

---

$$\left[ m\ddot{x} - \frac{x}{(x^2+y^2)^{\frac{3}{2}}} \quad m\ddot{y} - \frac{y}{(x^2+y^2)^{\frac{3}{2}}} \right]$$

Now we try it on a general central potential.

---

```
.../sage/section1.6.sage
703 U = Function(lambda x: function("U")(x))
704 show(Lagrange_equations(L_central_rectangular(m, U))(q)(t))
...
705
706
707
708
709
710
711
712
713
```

---

$$\left[ m\ddot{x} + \frac{x D_0(U)(\sqrt{x^2+y^2})}{\sqrt{x^2+y^2}} \quad m\ddot{y} + \frac{y D_0(U)(\sqrt{x^2+y^2})}{\sqrt{x^2+y^2}} \right]$$

### 1.6.4 Coordinate transformations

To get things straight: the function  $F$  is the transformation of the coordinates  $x'$  to  $x$ , i.e.,  $x = F(t, x')$ . The function  $C$  lifts the transformation  $F$  to the phase space, so it transforms  $\Gamma(q')$  to  $\Gamma(q)$ .

The result of  $\partial_1 F v$  is a vector, because  $v$  is a vector. We have to cast  $\partial_0 F$  into a vector to enable the summation of these two terms.

---

```
.../sage/utils1.6.sage
705 def F_to_C(F):
706     def f(local):
707         return up(
708             time(local),
709             F(local),
710             partial(F, 0)(local) + partial(F, 1)(local) * velocity(local),
711         )
712
713     return f
```

---

### 1.6.5 polar coordinates

---

```
.../sage/utils1.6.sage
714 def p_to_r(local):
715     r, phi = coordinate(local).list()
716     return column_matrix([r * cos(phi), r * sin(phi)])
```

---

We apply `F_to_C` and `p_to_r` to several examples, to test and to understand how they collaborate. We need to make the appropriate variables for the space in terms of  $r$  and  $\phi$ .

---

```
.../sage/section1.6.sage
717 r = literal_function("r")
718 phi = literal_function("phi")
719 q = column_path([r, phi])
720 show(p_to_r(Gamma(q)(t)))
```

---

$$\begin{bmatrix} \cos(\phi) r \\ r \sin(\phi) \end{bmatrix}$$

This is the derivative wrt  $t$ . As the transformation `p_to_r` does not depend explicitly on  $t$ , the result should be a column matrix of zeros.

---

```
.../sage/section1.6.sage
721 show((partial(p_to_r, 0)(Gamma(q)(t))))
```

---

$$\begin{bmatrix} 0 \\ 0 \end{bmatrix}$$

Next is the derivative wrt  $r$  and  $\phi$ .

---

```
.../sage/section1.6.sage
722 show((partial(p_to_r, 1)(Gamma(q)(t))))
```

---

$$\begin{bmatrix} \cos(\phi) & -r \sin(\phi) \\ \sin(\phi) & \cos(\phi) r \end{bmatrix}$$

---

```
.../sage/section1.6.sage
723 show(F_to_C(p_to_r)(Gamma(q)(t)))
```

---

$$\begin{aligned} t \\ \begin{bmatrix} \cos(\phi) r \\ r \sin(\phi) \end{bmatrix} \\ \begin{bmatrix} -r \sin(\phi) \dot{\phi} + \cos(\phi) \dot{r} \\ \cos(\phi) r \dot{\phi} + \sin(\phi) \dot{r} \end{bmatrix} \end{aligned}$$

We can see what happens for the Lagrangian for the central force in polar coordinates.

---

```
724 _____ ..../sage/utils1.6.sage _____
725 def L_central_polar(m, U):
726     def Lagrangian(local):
727         return compose(L_central_rectangular(m, U), F_to_C(p_to_r))(local)
728
729     return Lagrangian
```

---



---

```
729 _____ ..../sage/section1.6.sage _____
730 # show(L_central_polar(m, U)(Gamma(q)(t)))
731 show(L_central_polar(m, U)(Gamma(q)(t)).simplify_full())
```

---

$$\frac{1}{2}mr^2\dot{\phi}^2 + \frac{1}{2}mr^2 - U(\sqrt{r^2})$$

---

```
731 expr = Lagrange_equations(L_central_polar(m, U))(q)(t)
732 show(expr.simplify_full().expand())
```

---

$$\begin{bmatrix} -mr\dot{\phi}^2 + m\ddot{r} + \frac{rD_0(U)(\sqrt{r^2})}{\sqrt{r^2}} & mr^2\ddot{\phi} + 2mr\dot{\phi}\dot{r} \end{bmatrix}$$

### 1.6.6 Coriolis and centrifugal forces

---

```
733 _____ ..../sage/utils1.6.sage _____
734 def L_free_rectangular(m):
735     def Lagrangian(local):
736         v = velocity(local)
737         return 1 / 2 * m * square(v)
738
739     return Lagrangian
740
741 def L_free_polar(m):
742     def Lagrangian(local):
743         return L_free_rectangular(m)(F_to_C(p_to_r)(local))
744
745     return Lagrangian
746
747
748 def F(Omega):
749     def f(local):
750         t = time(local)
751         r, theta = coordinate(local).list()
752         return vector([r, theta + Omega * t])
753
754     return f
755
756 def L_rotating_polar(m, Omega):
```

```

758     def Lagrangian(local):
759         return L_free_polar(m)(F_to_C(F(Omega)))(local)
760
761     return Lagrangian
762
763
764
765     def r_to_p(local):
766         x, y = coordinate(local).list()
767         return column_matrix([sqrt(x * x + y * y), atan(y / x)])
768
769
770     def L_rotating_rectangular(m, Omega):
771         def Lagrangian(local):
772             return L_rotating_polar(m, Omega)(F_to_C(r_to_p))(local)
773
774     return Lagrangian


---


775     ..../sage/section1.6.sage
776     _ = var("Omega", domain="positive")
777     q_xy = column_path([literal_function("x"), literal_function("y")])
778     expr = L_rotating_rectangular(m, Omega)(Gamma(q_xy))(t).simplify_full()


---


778     ..../sage/section1.6.sage
    show(expr)


---



```

$$\frac{1}{2}\Omega^2 mx^2 + \frac{1}{2}\Omega^2 my^2 - \Omega my\dot{x} + \Omega mx\dot{y} + \frac{1}{2}m\ddot{x}^2 + \frac{1}{2}m\ddot{y}^2$$

The simplification of the Lagrange equations takes some time.

```

779     ..../sage/section1.6.sage
780     don't tangle
expr = Lagrange_equations(L_rotating_rectangular(m, Omega))(q)(t)
show(expr.simplify_full())


---



```

I edited the result a bit by hand.

$$-m\Omega^2 x - 2m\Omega\dot{y} + m\ddot{x}, -m\Omega^2 y + 2m\Omega\dot{x} + m\ddot{y}.$$

### 1.6.7 Constraints, a driven pendulum

Rather than implementation the formulas of the book at this place, we follow the idea they explain a bit later in the book: formulate a Lagrangian in practical coordinates, then formulate the problem in practical coordinates *for that problem*, and then use a coordinate transformation from the problem's coordinates to the Lagrangian coordinates.

For the driven pendulum, the Lagrangian is easiest to express in terms of  $x$  and  $y$  coordinates, while the pendulum needs an angle  $\theta$ . So, we need a transformation from

$\theta$  to  $x$  and  $y$ . Note that the function coordinate returns a  $(1 \times 1)$  column matrix which just contains  $\theta$ . So, we have to pick element  $(0,0)$ . Another point is that here  $ys$  needs to be evaluated at  $t$ ; in the other functions  $ys$  is just passed on as a function.

```
781     def dp_coordinates(l, ys):
782         "From theta to x, y coordinates."
783         def f(local):
784             t = time(local)
785             theta = coordinate(local)[0, 0]
786             return column_matrix([l * sin(theta), ys(t) - l * cos(theta)])
787
788     return f


---


789
790     def L_pend(m, l, g, ys):
791         def wrap_L_pend(local):
792             return L_uniform_acceleration(m, g)(
793                 F_to_C(dp_coordinates(l, ys))(local)
794             )
795
796         return wrap_L_pend


---


796     _ = var("l", domain="positive")
797
798     theta = column_path([literal_function("theta")])
799     ys = literal_function("y")
800
801     expr = L_pend(m, l, g, ys)(Gamma(theta)(t)).simplify_full()
802     show(expr)
```

$$\frac{1}{2} l^2 m \dot{\theta}^2 + lm \sin(\theta) \dot{\theta} \dot{y} + glm \cos(\theta) - gmy + \frac{1}{2} m \dot{y}^2$$

## 1.7 EVOLUTION OF DYNAMICAL STATE

### 1.7.1 Standard imports

```
803     load("utils1.6.sage")


---


804     load("utils1.7.sage")
805
806     var("t", domain=RR)


---


807     load("show_expression.sage")
```

### 1.7.2 Acceleration and state derivative

We build the functions `Lagrangian_to_acceleration` and `Lagrangian_to_state_derivative` in steps.

---

```
808     ..../sage/section1.7.sage
809     q = column_path([literal_function("x"), literal_function("y")])
810     local = Gamma(q)(t)
811     m, k = var("m k", domain="positive")
812     L = L_harmonic(m, k)
813     show(L(local))
```

---

$$-\frac{1}{2} (x^2 + y^2)k + \frac{1}{2} (\dot{x}^2 + \dot{y}^2)m$$

---

```
813     F = compose(transpose, partial(L, 1))
814     show(F(local))
815     P = partial(L, 2)
816     show((F - partial(P, 0))(local))
```

---

$$\begin{pmatrix} -kx \\ -ky \end{pmatrix}$$

$$\begin{pmatrix} -kx \\ -ky \end{pmatrix}$$

---

```
817     ..../sage/section1.7.sage
show((partial(P, 1) * velocity)(local))
```

---

$$\begin{pmatrix} 0 \\ 0 \end{pmatrix}$$

Convert to vector.

---

```
818     ..../sage/section1.7.sage
show((F - partial(P, 0) - partial(P, 1) * velocity)(local))
```

---

$$\begin{pmatrix} -kx \\ -ky \end{pmatrix}$$

---

```
819     ..../sage/utils1.7.sage
820     def Lagrangian_to_acceleration(L):
821         def f(local):
822             P = partial(L, 2)
823             F = compose(transpose, partial(L, 1))
824             M = (F - partial(P, 0)) - partial(P, 1) * velocity
825             return partial(P, 2)(local).solve_right(M(local))
826
return f
```

---

We apply this to the harmonic oscillator.

---

```
827   _____ ..../sage/section1.7.sage _____
show(Lagrangian_to_acceleration(L)(local))
```

---

$$\begin{pmatrix} -\frac{kx}{m} \\ -\frac{ky}{m} \end{pmatrix}$$

### 1.7.3 *Intermezzo, numerically integrating ODEs with Sagemath*

At a later stage, we want to numerically integrate the system of ODEs that result from the Lagrangian. This works a bit different from what I expected; here are two examples to see the problem.

Consider the system of DEs for the circle:  $\dot{x} = y$ ,  $\dot{y} = -x$ . This code implements the rhs:

---

```
828  _____ don't tangle _____
829  def de_rhs(x, y):
830      return [y, -x]
831
832  sol = desolve_odeint(de_rhs(x, y), [1, 0], srange(0, 100, 0.05), [x, y])
833  pp = list(zip(sol[:, 0], sol[:, 1]))
834  p = points(pp, color='blue', size=3)
835  p.save('circle.png')
```

---

However, if I replace the RHS of the DE by by constants,, I get an error that the integration variables are unknown.

---

```
836  _____ don't tangle _____
837  def de_rhs(x, y):
838      return [1, -1]
```

---

The solution is to replace the numbers by expressions.

---

```
838  _____ ..../sage/utils1.7.sage _____
839  def convert_to_expr(n):
840      return SR(n)
```

---

And then define the function of differentials like this.

---

```
840  _____ don't tangle _____
841  def de_rhs(x, y):
842      return [convert_to_expr(1), convert_to_expr(-1)]
```

---

Now things work as they should.

### 1.7.4 Continuing with the oscillator

The next function computes the state derivative of the Lagrangian. For the purpose of numerical integration, we cast the result of the derivative of  $dt/dt = 1$  to an expression, more specifically, by the above intermezzo we should set the derivative of  $t$  to `convert_to_expr(1)`.

```
..... ./sage/utils1.7.sage
842 def Lagrangian_to_state_derivative(L):
843     acceleration = Lagrangian_to_acceleration(L)
844     return lambda state: up(
845         convert_to_expr(1), velocity(state), acceleration(state)
846     )
..... ./sage/section1.7.sage
847 show(Lagrangian_to_state_derivative(L)(local))
..... ./sage/section1.7.sage
848 def harmonic_state_derivative(m, k):
849     return Lagrangian_to_state_derivative(L_harmonic(m, k))
..... ./sage/section1.7.sage
850 show(harmonic_state_derivative(m, k)(local))
..... ./sage/utils1.7.sage
851 def qv_to_state_path(q, v):
852     return lambda t: up(t, q(t), v(t))
..... ./sage/utils1.7.sage
853 def Lagrange_equations_first_order(L):
854     def f(q, v):
855         state_path = qv_to_state_path(q, v)
856         res = D(state_path)
857         res -= compose(Lagrangian_to_state_derivative(L), state_path)
858         return res
859
860     return f
```

---

```
861 _____ ./.sage/section1.7.sage _____
862 res = Lagrange_equations_first_order(L_harmonic(m, k))(
863     column_path([literal_function("x"), literal_function("y")]),
864     column_path([literal_function("v_x"), literal_function("v_y")]),
865 )
866 show(res(t))
```

---

$$\begin{pmatrix} 0 \\ -v_x + \dot{x} \\ -v_y + \dot{y} \\ \left( \begin{array}{c} \frac{kx}{m} + \dot{v}_x \\ \frac{ky}{m} + \dot{v}_y \end{array} \right) \end{pmatrix}$$

### 1.7.5 Numerical integration

For the numerical integrator we have to specify the variables that appear in the differential equations. For this purpose we use dummy vectors.

---

```
866 _____ ./.sage/utils1.7.sage _____
867 def make_dummy_vector(name, dim):
868     return column_matrix([var(f"{name}[{i}]", domain=RR) for i in range(dim)])
```

---

The state\_advancer needs an evolve function. We use the initial conditions ics to figure out the dimension of the coordinate space. Once we have the dimension, we construct a dummy up tuple with coordinate and velocity variables. The ode solver need plain lists; since space is an up tuple, the list method of Tuple can provide for this.

---

```
868 _____ ./.sage/utils1.7.sage _____
869 def evolve(state_derivative, ics, times):
870     dim = coordinate(ics).nrows()
871     coordinates = make_dummy_vector("q", dim)
872     velocities = make_dummy_vector("v", dim)
873     space = up(t, coordinates, velocities)
874     soln = desolve_odeint(
875         des=state_derivative(space).list(),
876         ics=ics.list(),
877         times=times,
878         dvars=space.list(),
879         atol=1e-13,
880     )
881     return soln
```

---

The state advancer integrates the orbit for a time T and starting at the initial conditions.

---

```
881 _____ ..../sage/utils1.7.sage _____
882 def state_advancer(state_derivative, ics, T):
883     init_time = time(ics)
884     times = [init_time, init_time + T]
885     soln = evolve(state_derivative, ics, times)
886     return soln[-1]
```

---

As a test, let's apply it to the one D harmonic oscillator.

---

```
886 state_advancer(
887     harmonic_state_derivative(m=2, k=1),
888     ics=up(0, column_matrix([1, 2]), column_matrix([3, 4])),
889     T=10,
890 )
```

---

array([10. , 3.71279102, 5.42061989, 1.61480284, 1.8189101 ])

These are (nearly) the same results as in the book.

### 1.7.6 The driven pendulum

Here is the driver for the pendulum.

---

```
891 _____ ..../sage/utils1.7.sage _____
892 def periodic_drive(amplitude, frequency, phase):
893     def f(t):
894         return amplitude * cos(frequency * t + phase)
895     return f
```

---

With this we make the Lagrangian.

---

```
896 _ = var("m l g A omega")
897
898 def L_periodically_driven_pendulum(m, l, g, A, omega):
899     ys = periodic_drive(A, omega, 0)
900
901     def Lagrangian(local):
902         return L_pend(m, l, g, ys)(local)
903
904     return Lagrangian
```

---



---

```
906 q = column_path([literal_function("theta")])
907 show(
908     L_periodically_driven_pendulum(m, l, g, A, omega)(
909         Gamma(q)(t)
910     ).simplify_full()
911 )
```

---

$$\frac{1}{2} A^2 m \omega^2 \sin(\omega t)^2 - A l m \omega \sin(\omega t) \sin(\theta) \dot{\theta} + \frac{1}{2} l^2 m \dot{\theta}^2 - A g m \cos(\omega t) + g l m \cos(\theta)$$

---

```

912      ..../sage/section1.7.sage
913  expr = Lagrange_equations(L_periodically_driven_pendulum(m, l, g, A, omega))(
914      q
915  )(t).simplify_full()
916  show(expr)

```

---

$$( l^2 m \ddot{\theta} - (A l m \omega^2 \cos(\omega t) - g l m) \sin(\theta) )$$

---

```

916      ..../sage/section1.7.sage
917  show(
918      Lagrangian_to_acceleration(
919          L_periodically_driven_pendulum(m, l, g, A, omega)
920      )(Gamma(q)(t)).simplify_full()
921  )

```

---

$$\left( \frac{(A\omega^2 \cos(\omega t) - g) \sin(\theta)}{l} \right)$$

---

```

921 def pend_state_derivative(m, l, g, A, omega):
922     return Lagrangian_to_state_derivative(
923         L_periodically_driven_pendulum(m, l, g, A, omega)
924     )

```

---



---

```

925 expr = pend_state_derivative(m, l, g, A, omega)(Gamma(q)(t))
926 show(time(expr))
927 show(coordinate(expr).simplify_full())
928 show(velocity(expr).simplify_full())

```

---

1

$$(\dot{\theta})$$

$$\left( \frac{(A\omega^2 \cos(\omega t) - g) \sin(\theta)}{l} \right)$$

---

```

929 def principal_value(cut_point):
930     def f(x):
931         return (x + cut_point) % (2 * np.pi) - cut_point
932
933     return f

```

---

---

```

934 _____ ..../sage/section1.7.sage _____
935 def plot_driven_pendulum(A, T, step_size=0.01):
936     times = sage(range(0, T, step_size, include_endpoint=True))
937     soln = evolve(
938         pend_state_derivative(m=1, l=1, g=9.8, A=A, omega=2 * sqrt(9.8)),
939         ics=up(0, column_matrix([1]), column_matrix([0])),
940         times=times,
941     )
942     thetas = soln[:, 1]
943     pp = list(zip(times, thetas))
944     p = points(pp, color='blue', size=3)
945     p.save(f'..//figures/driven_pendulum_{A:.2f}.png')
946
947     thetas = principal_value(np.pi)(thetas)
948     pp = list(zip(times, thetas))
949     p = points(pp, color='blue', size=3)
950     p.save(f'..//figures/driven_pendulum_{A:.2f}_principal_value.png')
951
952     thetadots = soln[:, 2]
953     pp = list(zip(thetas, thetadots))
954     p = points(pp, color='blue', size=3)
955     p.save(f'..//figures/driven_pendulum_{A:.2f}_trajectory.png')

```

---

So now we make the plot.

---

```

956 _____ ..../sage/section1.7.sage _____
plot_driven_pendulum(A=0.1, T=100, step_size=0.005)

```

---

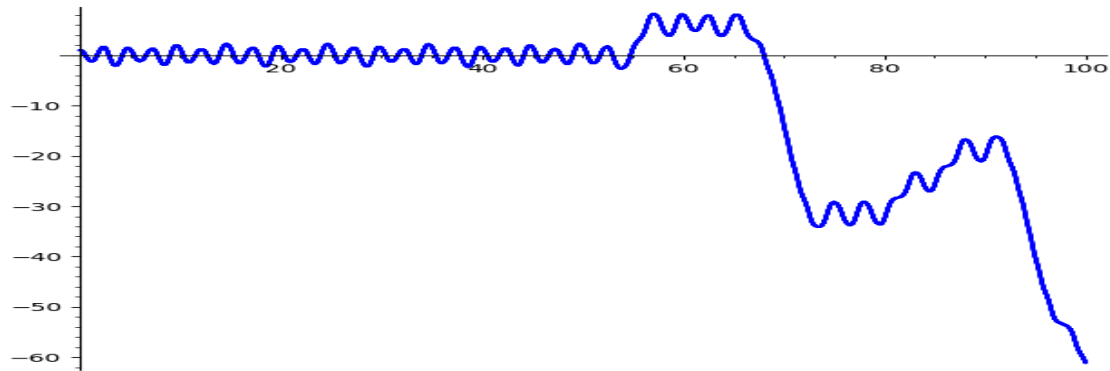
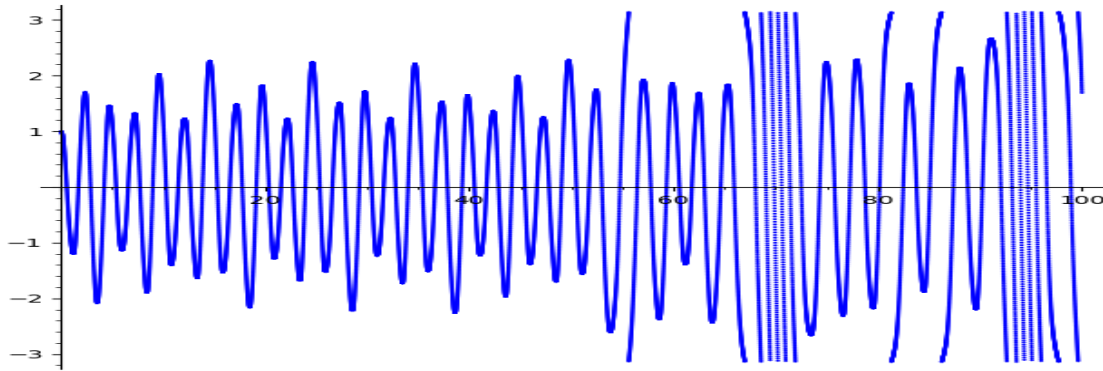
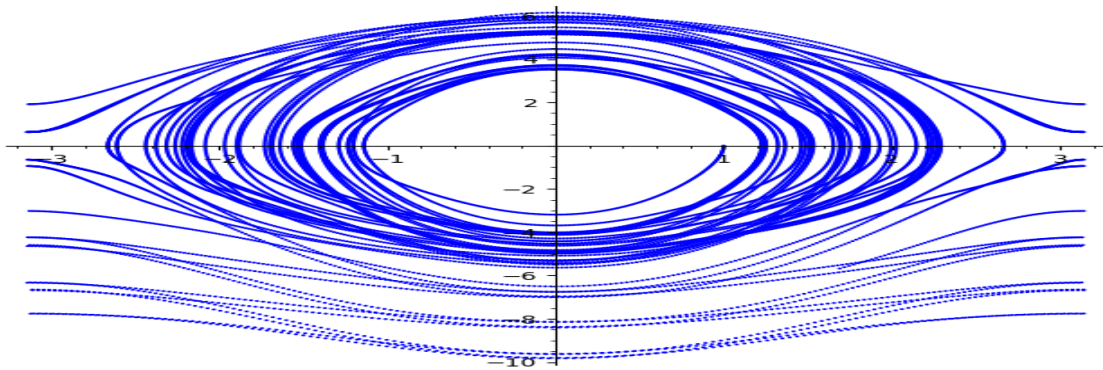


Figure 1.2: The angle of the vertically driven pendulum as a function of time. Obviously, around  $t = 80$ , the pendulum makes a few revolutions, and then starts to wobble again.

Figure 1.3: The angle on  $(-\pi, \pi]$ .Figure 1.4: The trajectory of  $\theta$  and  $\dot{\theta}$ .

## 1.8 CONSERVED QUANTITIES

### 1.8.1 Standard imports

```

957   _____ .. / sage / utils1.8.sage _____
load( "utils1.6.sage" )

958   _____ .. / sage / section1.8.sage _____
load( "utils1.8.sage" )
959
960 var( "t" , domain=RR)
961
         _____ don't tangle _____
load( "show_expression.sage" )

```

### 1.8.2 1.8.2 Energy Conservation

From the Lagrangian we can construct the energy function. Note that we should cast  $P = \partial_2 L$  to a vector so that  $P * v$  becomes a number instead of a  $1 \times 1$  matrix. As we use the Lagrangian in functional arithmetic, we convert  $L$  into a Function.

---

```
962 def Lagrangian_to_energy(L):
963     P = partial(L, 2)
964     LL = Function(lambda local: L(local))
965     return lambda local: (P * velocity - LL)(local)
```

---

### 1.8.3 Central Forces in Three Dimensions

Instead of building the kinetic energy in spherical coordinates, as in Section 1.8.3 of the book, I am going to use the ideas that have been expounded book in earlier sections: define the Lagrangian in convenient coordinates, and then use a coordinate transform to obtain it in coordinates that show the symmetries of the system.

---

```
966 q = column_path(
967     [
968         literal_function("r"),
969         literal_function("theta"),
970         literal_function("phi"),
971     ]
972 )
```

---

Next the transformation from spherical to 3D rectangular coordinates.

---

```
973 def s_to_r(spherical_state):
974     r, theta, phi = coordinate(spherical_state).list()
975     return vector(
976         [r * sin(theta) * cos(phi), r * sin(theta) * sin(phi), r * cos(theta)])
977 )
```

---

For example, here is are the velocities expressed in spherical coordinates.

---

```
978 show(velocity(F_to_C(s_to_r))(Gamma(q)(t))).simplify_full()
```

---

$$\begin{bmatrix} \cos(\phi)\cos(\theta)r\dot{\theta} - (r\sin(\phi)\dot{\phi} - \cos(\phi)\dot{r})\sin(\theta) \\ \cos(\theta)r\sin(\phi)\dot{\theta} + (\cos(\phi)r\dot{\phi} + \sin(\phi)\dot{r})\sin(\theta) \\ -r\sin(\theta)\dot{\theta} + \cos(\theta)\dot{r} \end{bmatrix}$$

Now we are ready to check the code examples of the book.

---

```
979 V = Function(lambda r: function("V")(r))
980
981 def L_3D_central(m, V):
982     def Lagrangian(local):
983         return L_central_rectangular(m, V)(F_to_C(s_to_r)(local))
984
985     return Lagrangian
```

---

---

```

986 _____ ..../sage/section1.8.sage _____
986 show(partial(L_3D_central(m, V), 1)(Gamma(q)(t)).simplify_full())
_____  


$$\left[ -\frac{rD_0(V)\sqrt{r^2} - (mr\sin(\theta)^2\dot{\phi}^2 + mr\dot{\theta}^2)\sqrt{r^2}}{\sqrt{r^2}} \quad m\cos(\theta)r^2\sin(\theta)\dot{\phi}^2 \quad 0 \right]$$

987 _____ ..../sage/section1.8.sage _____
987 show(partial(L_3D_central(m, V), 2)(Gamma(q)(t)).simplify_full())
_____  


$$\left[ m\dot{r} \quad mr^2\dot{\theta} \quad mr^2\sin(\theta)^2\dot{\phi} \right]$$

_____  

988 def ang_mom_z(m):
989     def f(rectangular_state):
990         xyx = vector(coordinate(rectangular_state))
991         v = vector(velocity(rectangular_state))
992         return xyx.cross_product(m * v)[2]
993
994     return f
995
996
997 show(compose(ang_mom_z(m), F_to_C(s_to_r))(Gamma(q)(t)).simplify_full())

```

---

$$mr^2\sin(\theta)^2\dot{\phi}$$

This is the check that  $E = T + V$ .

---

```

998 _____ ..../sage/section1.8.sage _____
998 show(Lagrangian_to_energy(L_3D_central(m, V))(Gamma(q)(t)).simplify_full())
_____  


$$\left[ \frac{1}{2}mr^2\sin(\theta)^2\dot{\phi}^2 + \frac{1}{2}mr^2\dot{\theta}^2 + \frac{1}{2}m\dot{r}^2 + V(\sqrt{r^2}) \right]$$


```

---

#### 1.8.4 The Restricted Three-Body Problem

I decompose the potential energy function into smaller functions; I find the implementation in the book somewhat heavy.

---

```

999 var("G M0 M1 a", domain="positive")
1000
1001
1002 def distance(x, y):
1003     return sqrt(square(x - y))
1004
1005
1006 def angular_freq(M0, M1, a):

```

---

```

1007     return sqrt(G * (M0 + M1) / a ^ 3)
1008
1009
1010 def V(a, M0, M1, m):
1011     Omega = angular_freq(M0, M1, a)
1012     a0, a1 = M1 / (M0 + M1) * a, M0 / (M0 + M1) * a
1013
1014     def f(t, origin):
1015         pos0 = -a0 * column_matrix([cos(Omega * t), sin(Omega * t)])
1016         pos1 = a1 * column_matrix([cos(Omega * t), sin(Omega * t)])
1017         r0 = distance(origin, pos0)
1018         r1 = distance(origin, pos1)
1019         return -G * m * (M0 / r0 + M1 / r1)
1020
1021     return f
1022
1023 def L0(m, V):
1024     def f(local):
1025         t, q, v = time(local), coordinate(local), velocity(local)
1026         return 1 / 2 * m * square(v) - V(t, q)
1027
1028 return f

```

---

For the computer it's easy to compute the energy, but the formula is pretty long.

```

----- ./sage/section1.8.sage -----
1029 q = column_path([literal_function("x"), literal_function("y")])
1030 expr = (sqrt(G*M0 + G*M1)*t) / a^(3/2)
1031 A = var('A')
1032
1033 show(
1034     Lagrangian_to_energy(L0(m, V(a, M0, M1, m)))(Gamma(q)(t))
1035     .simplify_full()
1036     .expand()
1037     .subs({expr: A})
1038 )

```

---

$$\left[ -\frac{\sqrt{M_0^2+2 M_0 M_1+M_1^2} G M_0 m}{\sqrt{2 M_0 M_1 a \cos(A) x+2 M_1^2 a \cos(A) x+2 M_0 M_1 a \sin(A) y+2 M_1^2 a \sin(A) y+M_1^2 a^2+M_0^2 x^2+2 M_0 M_1 x^2+M_1^2 x^2+M_0^2 y^2+2 M_0 M_1 y^2+M_1^2 y^2}} \right]$$

I skip the rest of the code of this part as it is just copy work from the mathematical formulas.

### 1.8.5 Noether's theorem

We need to rotate around a given axis in 3D space. ChatGPT gave me the code right away.

```

----- ./sage/utils1.8.sage -----
1039 def rotation_matrix(axis, theta):

```

```

1040 """
1041 Return the 3x3 rotation matrix for a rotation of angle theta (in radians)
1042 about the given axis. The axis is specified as an iterable of 3 numbers.
1043 """
1044 # Convert the axis to a normalized vector
1045 axis = vector(axis).normalized()
1046 x, y, z = axis
1047 c = cos(theta)
1048 s = sin(theta)
1049 t = 1 - c # common factor
1050
1051 # Construct the rotation matrix using Rodrigues' formula
1052 R = matrix(
1053     [
1054         [c + x**2 * t, x * y * t - z * s, x * z * t + y * s],
1055         [y * x * t + z * s, c + y**2 * t, y * z * t - x * s],
1056         [z * x * t - y * s, z * y * t + x * s, c + z**2 * t],
1057     ]
1058 )
1059


---



```

```

.../sage/section1.8.sage
1060 def F_tilde(angle_x, angle_y, angle_z):
1061     def f(local):
1062         return (
1063             rotation_matrix([1, 0, 0], angle_x)
1064             * rotation_matrix([0, 1, 0], angle_y)
1065             * rotation_matrix([0, 0, 1], angle_z)
1066             * coordinate(local)
1067         )
1068
1069     return f


---



```

```

.../sage/section1.8.sage
1070 q = column_path(
1071     [literal_function("x"), literal_function("y"), literal_function("z")]
1072 )


---



```

Let's see what we get when we exercise a rotation of  $s$  radians round the  $x$  axis.

```

.../sage/section1.8.sage
1073 def Rx(s):
1074     return lambda local: F_tilde(s, 0, 0)(local)
1075
1076
1077 s, u, v = var("s u v")
1078 latex.matrix_delimiters(left='[', right=']')
1079 latex.matrix_column_alignment("c")
1080 show(Rx(s)(Gamma(q)(t)))
1081 show(diff(Rx(s)(Gamma(q)(t)), s)(s=0))


---



```

$$\begin{bmatrix} x \\ \cos(s)y - \sin(s)z \\ \sin(s)y + \cos(s)z \end{bmatrix}$$

$$\begin{bmatrix} 0 \\ -z \\ y \end{bmatrix}$$

And now we check the result of the book. The computation of  $D F_{\tilde{t}}$  is somewhat complicated. Observe that  $F_{\tilde{t}}$  is a function of the rotation angles, and returns a function that takes `local` as argument. Now we want to differentiate  $F_{\tilde{t}}$  with respect to the angles, so these are the variables we need to provide to the Jacobian. For this reason, we bind the result of  $F_{\tilde{t}}$  to `local`, and use a lambda function to provide the angles as the variables. This gives us  $F_{\tilde{t}}$  (note that I drop the underscore in this name). There is one further point:  $F_{\tilde{t}}$  expects three angles, while the Jacobian provides the list  $[s, u, v]$  as the argument to  $F_{\tilde{t}}$ . Therefore we unpack the argument `x` of the lambda function to convert the list  $[s, u, v]$  into three separate arguments. The last step is to fill in  $s = u = v = 0$ .

Note that we differentiate wrt  $s, u, v$  and not wrt  $t$ . In itself, using  $t$  would not be a problem, but since we pass  $\Gamma(q)(t)$  to  $F_{\tilde{t}}$ , the function depends also on  $t$  via the path  $t \rightarrow \Gamma(q, t)$  which we should avoid.

As for the result, I don't see why my result differs by a minus sign from the result in the book.

---

```

1082 U = Function(lambda r: function("U")(r))
1083
1084
1085 def the_Noether_integral(local):
1086     L = L_central_rectangular(m, U)
1087     Ftilde = lambda x: F_tilde(*x)(local)
1088     DF0 = Jacobian(Ftilde)([s, u, v], [s, u, v])(s=0, u=0, v=0)
1089     return partial(L, 2)(local) * DF0


---


1090 show(the_Noether_integral(Gamma(q)(t)).simplify_full())

```

---

$$\begin{bmatrix} -mz\dot{y} + my\dot{z} & mz\dot{x} - mx\dot{z} & -my\dot{x} + mx\dot{y} \end{bmatrix}$$

## 1.9 ABSTRACTION OF PATH FUNCTIONS

I found this section difficult to understand, so I work in small steps to the final result, and include checks to see what goes on.

### 1.9.1 Standard imports

```
1091      ..... /sage/utils1.9.sage
load("utils1.6.sage")  

1092      ..... /sage/section1.9.sage
load("utils1.9.sage")  

1093  

1094 var("t", domain=RR)  

1095      ..... don't tangle
load("show_expression.sage")
```

### 1.9.2 Understanding *F\_to\_C*

The Scheme code starts with defining `Gamma_bar` in terms of `f_bar` and `osculating_path`. We build `f_bar` first and apply it to the example in which polar coordinates are converted to rectilinear coordinates.

Next, let's spell out the arguments of all functions to see how everything works together. A literal function maps time  $t$  to some part of the space, often to a coordinate,  $x$  say.

```
1096      ..... /sage/section1.9.sage
r, theta = literal_function("r"), literal_function("theta")
show(r)  

<__main__.Function object at 0x752ed4eb27a0>
```

So, `r` is a Function. We can evaluate `r` at  $t$ . I pass `simplify=False` to show to *not* suppress the dependence on  $t$ .

```
1098      ..... /sage/section1.9.sage
show((r(t), theta(t)), simplify=False)  

(r(t), theta(t))
```

A `column_path` takes literal functions as arguments and returns a coordinate path. Hence, it is a function of  $t$  and returns  $q(t)$ . (I use the notation of the code examples of the book such as `q_prime` so that I can copy the examples into the functions I build later.)

```
1099      ..... /sage/section1.9.sage
q_prime = column_path([r, theta])
show(q_prime(t), simplify=False)
```

$$\begin{bmatrix} r(t) \\ \theta(t) \end{bmatrix}$$

The function  $\Gamma$  takes a coordinate path  $q$  (which is a function of time) as input, and returns a function of  $t$  that maps to a local up tuple  $l$ :

$$\Gamma[q] : t \rightarrow l = (t, q(t), v(t), \dots).$$

---

```
1101      show(Gamma(q_prime))
```

---

```
<function Gamma.<locals>.<lambda> at 0x752ed4ba0cc0>
```

Indeed, `Gamma` is a function, and has to be applied to some argument to result into a value. In fact, when  $\Gamma(q)$  is applied to  $t$ , we get the local up tuple  $l$ . Observe, that a local tuple is *not* a functions of time, by that I mean, a local is not a Python function of time, and therefore does not take any further arguments.

---

```
1102      show(Gamma(q_prime)(t), simplify=False)
```

---

$$\begin{bmatrix} t \\ r(t) \\ \theta(t) \\ \frac{\partial}{\partial t} r(t) \\ \frac{\partial}{\partial t} \theta(t) \end{bmatrix}$$

The coordinate transformation  $F$  in the example that transforms polar coordinates to rectilinear coordinates is `p_to_r`. This transform  $F$  maps a local tuple  $l$  to coordinates  $q(t)$ . Therefore, we can apply  $F$  to  $\Gamma[q](t)$ , and use composition like this:

$$F(\Gamma[q](t)) = (F \circ \Gamma[q])(t).$$

Observe that  $F \circ \Gamma[q]$  is a function of  $t$ .

---

```
1103      F = p_to_r
1104      show(compose(F, Gamma(q_prime))(t), simplify=False)
```

---

$$\begin{bmatrix} \cos(\theta(t))r(t) \\ r(t)\sin(\theta(t)) \end{bmatrix}$$

Since  $F \circ \Gamma[q]$  is a function of  $t$  to a coordinate path  $q(t)$ , this function has the same ‘protocol’ as a coordinate path function. We can therefore apply  $\Gamma$  to the composite function  $F \circ \Gamma[q]$  to obtain a function that maps  $t$  to a local tuple in the transformed space.

$$Q : t \rightarrow \Gamma[F \circ \Gamma[q]](t).$$

---

```
1105 Q = lambda t: compose(p_to_r, Gamma(q_prime))(t)
1106 show(Gamma(Q)(t), simplify=False)
```

---

$$\begin{bmatrix} t \\ \cos(\theta(t))r(t) \\ r(t)\sin(\theta(t)) \\ -r(t)\sin(\theta(t))\frac{\partial}{\partial t}\theta(t) + \cos(\theta(t))\frac{\partial}{\partial t}r(t) \\ \cos(\theta(t))r(t)\frac{\partial}{\partial t}\theta(t) + \sin(\theta(t))\frac{\partial}{\partial t}r(t) \end{bmatrix}$$

Now that we have analyzed all steps, we can make `f_bar`.

---

```
1107 def f_bar(q_prime):
1108     q = lambda t: compose(F, Gamma(q_prime))(t)
1109     return lambda t: Gamma(q)(t)
```

---

Here is the check. I suppress the dependence on  $t$  again to keep the result easier to read.

---

```
1110 show(f_bar(q_prime)(t))
```

---

$$\begin{bmatrix} t \\ \cos(\theta)r \\ r\sin(\theta) \\ -r\sin(\theta)\dot{\theta} + \cos(\theta)\dot{r} \\ \cos(\theta)r\dot{\theta} + \sin(\theta)\dot{r} \end{bmatrix}$$

The second function to build is `osculating_path`. This is the Taylor series of the book in which a local tuple is mapped to coordinate space:

$$O(t, q, v, a, \dots)(\cdot) = q + v(\cdot - t) + a/2(\cdot - t)^2 + \dots$$

I write  $\cdot$  instead of  $t'$  to make explicit that  $O(l)$  is still a function of  $t'$  in this case.

Clearly, the RHS is a sum of vectors all of which have the same dimension as the space of coordinates.

Rather than computing  $dt^n$  as  $(t - t')^n$ , and  $n!$  for each  $n$ , I compute these values recursively. The implementation assumes that the local tuple  $\Gamma[q](t)$  contains at least the elements  $t$  and  $q$ , that is  $\Gamma[q](t) = (t, q, \dots)$ . This local tuple has length 2; the local tuple  $l = (t, q, v)$  has length 3.

---

```
1111 def osculating_path(local):
1112     t = time(local)
1113     q = coordinate(local)
```

---

```

1114
1115     def wrapper(t_prime):
1116         res = q
1117         dt = 1
1118         factorial = 1
1119         for k in range(2, len(local)):
1120             factorial *= k
1121             dt *= t_prime - t
1122             res += local[k] * dt / factorial
1123         return res
1124
1125     return wrapper

```

---

Here is an example.

```

----- ./sage/section1.9.sage -----
1126 t_prime = var("tt", domain="positive", latex_name="t' ")
1127 q = column_path([literal_function("r"), literal_function("theta")])
1128 local = Gamma(q)(t)
1129 show(osculating_path(local)(t_prime))

```

---

$$\begin{bmatrix} -\frac{1}{2}(t-t')\dot{r} + r \\ -\frac{1}{2}(t-t')\dot{\theta} + \theta \end{bmatrix}$$

With the above pieces we can finally build `Gamma_bar`.

```

----- ./sage/utils1.9.sage -----
1130 def Gamma_bar(f_bar):
1131     def wrapped(local):
1132         t = time(local)
1133         q_prime = osculating_path(local)
1134         return f_bar(q_prime)(t)
1135
1136     return wrapped

```

---

```

----- ./sage/section1.9.sage -----
1137 show(Gamma_bar(f_bar)(local))

```

---

$$\begin{bmatrix} t \\ \cos(\theta)r \\ r\sin(\theta) \\ -r\sin(\theta)\dot{\theta} + \cos(\theta)\dot{r} \\ \cos(\theta)r\dot{\theta} + \sin(\theta)\dot{r} \end{bmatrix}$$

We can use `Gamma_bar` in to produce the transformation for polar to rectilinear coordinates.

---

```

1138 ..../sage/utils1.9.sage
1139 def F_to_C(F):
1140     def C(local):
1141         n = len(local)
1142
1143         def f_bar(q_prime):
1144             q = lambda t: compose(F, Gamma(q_prime))(t)
1145             return lambda t: Gamma(q, n)(t)
1146
1147         return Gamma_bar(f_bar)(local)
1148
1149     return C

```

---

```

1149 ..../sage/section1.9.sage
show(F_to_C(p_to_r)(local))

```

---

$$\begin{bmatrix} t \\ \cos(\theta) r \\ r \sin(\theta) \\ -r \sin(\theta) \dot{\theta} + \cos(\theta) \dot{r} \\ \cos(\theta) r \dot{\theta} + \sin(\theta) \dot{r} \end{bmatrix}$$

Here is the total time derivative.

---

```

1150 ..../sage/utils1.9.sage
@Func
1151 def Dt(F):
1152     def DtF(local):
1153         n = len(local)
1154
1155         def DF_on_path(q):
1156             return D(lambda t: F(Gamma(q, n - 1))(t))
1157
1158         return Gamma_bar(DF_on_path)(local)
1159
1160     return lambda state: DtF(local)

```

---

### 1.9.3 Lagrange equations at a moment

---

```

1161 ..../sage/utils1.9.sage
def Euler_Lagrange_operator(L):
    return lambda local: (Dt(partial(L, 2)) - partial(L, 1))(local)

```

---

To apply this operator to a local tuple, we need to include the acceleration.

---

```

1163 ..../sage/section1.9.sage
1164 q = column_path([literal_function("x")])
1165 local = Gamma(q, 4)(t)
1166 show(local)

```

---

$$\begin{bmatrix} t \\ x \\ \dot{x} \\ \ddot{x} \end{bmatrix}$$

---

.../sage/section1.9.sage

---

```
1166 m, k = var("m k", domain="positive")
1167 L = L_harmonic(m, k)
1168 show(Euler_Lagrange_operator(L)(local))
```

---

$$\begin{bmatrix} kx + m\ddot{x} \end{bmatrix}$$

# 2

CHAPTER 2: SKIPPED

---

## CHAPTER 3

---

### 3.1 HAMILTON'S EQUATIONS

#### 3.1.1 Standard imports

```
1169      ..... /sage/utils3.1.sage
load("utils1.6.sage")  
  

1170      ..... /sage/section3.1.sage
load("utils3.1.sage")  
  

1171      t = var("t", domain="real")  
  

1173      ..... don't tangle
load("show_expression.sage")  
  


```

---

#### 3.1.2 Computing Hamilton's equations

The code in Section 3.1 of the book starts with the following function.

```
..... /sage/utils3.1.sage
1174 def Hamilton_equations(Hamiltonian):
1175     def f(q, p):
1176         state_path = qp_to_H_state_path(q, p)
1177         return D(state_path) - compose(
1178             Hamiltonian_to_state_derivative(Hamiltonian), state_path
1179         )
1180
1181     return f  
  


```

---

This needs the next function.

```
..... /sage/utils3.1.sage
1182 def qp_to_H_state_path(q, p):
1183     def f(t):
1184         return up(t, q(t), p(t))
1185
1186     return f  
  


```

---

Here  $p$  is a function that maps  $t$  to a momentum vector. Such vectors are represented as lying vectors (or down tuples in the book). To implement this, `row_path` takes a list and returns a function that maps time to the transpose of a column path. In passing we

define `row_matrix` as the transpose of `column_matrix`, which is the function provided by Sagemath, and a `transpose` function.

---

```
1187 def transpose(M):
1188     return M.T
1189
1190
1191 def row_path(lst):
1192     return lambda t: transpose(column_path(lst)(t))
1193
1194
1195 def row_matrix(lst):
1196     return transpose(column_matrix(lst))
```

---

Let's try what we built.

---

```
1197 q = column_path([literal_function("q_x"), literal_function("q_y")])
1198 p = row_path([literal_function("p_x"), literal_function("p_y")])
1199
1200
1201 show(p(t))
```

---

$$\begin{bmatrix} p_x & p_y \end{bmatrix}$$

---

```
1200 H_state = qp_to_H_state_path(q, p)(t)
1201 show(H_state)
```

---

$$\begin{bmatrix} t \\ \begin{bmatrix} q_x \\ q_y \end{bmatrix} \\ \begin{bmatrix} p_x & p_y \end{bmatrix} \end{bmatrix}$$

The next function on which `Hamiltonian_equations` depends is `Hamiltonian_to_state_derivative`. The book prints the system of differential equations as a column vector. Therefore we transpose  $\partial_2 H$ .

---

```
1202 def Hamiltonian_to_state_derivative(Hamiltonian):
1203     def f(H_state):
1204         return up(
1205             SR(1),
1206             partial(Hamiltonian, 2)(H_state).T,
1207             -partial(Hamiltonian, 1)(H_state),
1208         )
1209
1210     return f
```

---

Here is an example with `H_rectangular`. For some reason, the book takes just the first and second component of `q`, i.e. (`req q 0`) and (`ref q 1`), to pass to the potential, but the general formula works just as well.

---

```
1211      var("m")
1212
1213  def H_rectangular(m, V):
1214      def f(state):
1215          q, p = coordinate(state), momentum(state)
1216          return square(p) / 2 / m + V(q)
1217
1218      return f
```

---

For this to work, we need a momentum projection operator. It's the same as the velocity projection.

---

```
1219  momentum = Function(lambda H_state: H_state[2])
```

---

Recall, to use symbolic functions in differentiation, the symbolic function requires an unpacked list of arguments.

---

```
1220  V = Function(lambda x: function("V")(*x.list()))
```

---

This is the Hamiltonian.

---

```
1221  H = H_rectangular
1222  show(H(m, V)(H_state))
```

---

$$\frac{p_x^2 + p_y^2}{2m} + V(q_x, q_y)$$

Partial derivatives work.

---

```
1223  show(partial(H(m, V), 1)(H_state))
```

---

$$[ D_0(V)(q_x, q_y) \quad D_1(V)(q_x, q_y) ]$$

---

```
1224  show(Hamiltonian_to_state_derivative(H(m, V))(H_state))
```

---

$$\begin{bmatrix} 1 \\ \begin{bmatrix} \frac{p_x}{m} & \frac{p_y}{m} \end{bmatrix} \\ \begin{bmatrix} -D_0(V)(q_x, q_y) & -D_1(V)(q_x, q_y) \end{bmatrix} \end{bmatrix}$$

---

```
1225 ..../sage/section3.1.sage
show(Hamilton_equations(H(m, V))(q, p)(t))


---



$$\begin{bmatrix} 0 \\ -\frac{p_x}{m} + \dot{q}_x \\ -\frac{p_y}{m} + \dot{q}_y \\ D_0(V)(q_x, q_y) + \dot{p}_x \quad D_1(V)(q_x, q_y) + \dot{p}_y \end{bmatrix}$$

```

### 3.1.3 The Legendre Transformation

To understand the code of the book, observe the following.

$$\begin{aligned} F(v) &= 1/2v^T M v + b^t v + c, \\ \partial_v F(v) &= M v + b, \\ \partial_v F(0) &= b, \\ \partial_v^2 F(v) &= M. \end{aligned}$$

Clearly,  $\partial_v F$  is the gradient, and  $\partial_v^2 F$  is the Hessian. Observe that under the operation of the gradient, the vector  $b$  changes shape: from  $b^t$  to  $b$ .

In the code, the argument  $w$  corresponds to a moment, hence is a lying vector. We need some dummy symbols with respect to which to differentiate, and then we set the dummy variables to  $o$  in the gradient and the Hessian. For this second step, Sagemath uses substitution with a dictionary when multiple arguments are involved, which is the case here because  $w$  is a vector. So, by making a `zeros` dictionary that maps symbols to  $o$ , we can use the keys of `zeros` as the dummy symbols, and then use `zeros` itself in the substitution. Then, to solve for  $v$  such that  $Mv = w^t - b$ , the lying vector  $w$  has to be transposed.

---

```
1226 ..../sage/utils3.1.sage
1227 def Legendre_transform(F):
1228     def G(w):
1229         zeros = {var(f"v_{i}"): 0 for i in range(w.ncols())}
1230         b = gradient(F)(list(zeros.keys())).subs(zeros)
1231         M = Hessian(F)(list(zeros.keys())).subs(zeros)
1232         v = M.solve_right(w.T - b)
1233         return w * v - F(v)
1234


---



```

Now we are equipped to convert a Lagrangian into a Hamiltonian.

---

```
1235 ..../sage/utils3.1.sage
1236 def Lagrangian_to_Hamiltonian(Lagrangian):
1237     def f(H_state):
1238         t = time(H_state)
```

```

1238     q = coordinate(H_state)
1239     p = momentum(H_state)
1240
1241     def L(qdot):
1242         return Lagrangian(up(t, q, qdot))
1243
1244     return Legendre_transform(L)(p)
1245
1246     return f

```

---

```

1247     ..... ./sage/section3.1.sage
1248 res = Lagrangian_to_Hamiltonian(L_central_rectangular(m, V))(H_state)
1249 show(res)

```

---

$$\left[ -\frac{1}{2}m\left(\frac{p_x^2}{m^2} + \frac{p_y^2}{m^2}\right) + \frac{p_x^2}{m} + \frac{p_y^2}{m} + V\left(\sqrt{q_x^2 + q_y^2}\right) \right]$$

```

1249 ..... ./sage/section3.1.sage
show(res.simplify_full())

```

---

$$\left[ \frac{2mV\left(\sqrt{q_x^2+q_y^2}\right)+p_x^2+p_y^2}{2m} \right]$$

```

1250 ..... ./sage/section3.1.sage
var("m g l")
1251 q = column_path([literal_function("theta")])
1252 p = row_path([literal_function("p")])

```

---

Here is exercise 3.1.

```

1253 ..... ./sage/section3.1.sage
# space = make_named_space(["\\theta"])
1254 H_state = qp_to_H_state_path(q, p)(t)
1255 show(Lagrangian_to_Hamiltonian(L_planar_pendulum(m, g, l))(H_state))

```

---

$$\left[ -g l m (\cos(\theta) - 1) + \frac{p^2}{2l^2 m} \right]$$

```

1256 ..... ./sage/section3.1.sage
q = column_path([literal_function("q_x"), literal_function("q_y")])
1257 p = row_path([literal_function("p_x"), literal_function("p_y")])
1258 H_state = qp_to_H_state_path(q, p)(t)
1259 show(Lagrangian_to_Hamiltonian(L_Henon_Heiles(m))(H_state))

```

---

$$\left[ q_x^2 q_y - \frac{1}{3} q_y^3 + \frac{1}{2} p_x^2 + \frac{1}{2} p_y^2 + \frac{1}{2} q_x^2 + \frac{1}{2} q_y^2 \right]$$

---

```

1260 _____ .. / sage / section3.1.sage _____
1261 def L_sphere(m, R):
1262     def Lagrangian(local):
1263         theta, phi = coordinate(local).list()
1264         thetadot, phidot = velocity(local).list()
1265         return 1 / 2 * m * R ^ 2 * (
1266             square(thetadot) + square(phidot * sin(theta)))
1267
1268     return Lagrangian
1269
1270
1271 var("R", domain="positive")

```

---



---

```

1272 _____ .. / sage / section3.1.sage _____
1273 q = column_path([literal_function("theta"), literal_function("phi")])
1274 p = row_path([literal_function("p_x"), literal_function("p_y")])
1275 H_state = qp_to_H_state_path(q, p)(t)
1276 show(Lagrangian_to_Hamiltonian(L_sphere(m, R))(H_state).simplify_full())

```

---

$$\left[ \frac{p_x^2 \sin(\theta)^2 + p_y^2}{2 R^2 m \sin(\theta)^2} \right]$$

## 3.2 POISSON BRACKETS

### 3.2.1 *The standard imports.*

### 3.2.2 *Standard imports*

---

```

1276 load("utils3.1.sage")

```

---

```

1277 load("utils3.2.sage")
1278 t = var("t", domain="real")

```

---

```

1280 load("show_expression.sage")

```

---

### 3.2.3 *The Poisson Bracket*

This is the Poisson bracket.

---

```

1281 @Func

```

---

```

1282 def Poisson_bracket(F, G):
1283     def f(state):
1284         left = (partial(F, 1) * compose(transpose, partial(G, 2)))(state)
1285         right = (partial(F, 2) * compose(transpose, partial(G, 1)))(state)
1286         return (left - right).simplify_full()
1287
1288     return f

```

---

We can make general state functions like so.

```

1289 @Func
1290 def state_function(name):
1291     return lambda H_state: function(name)(
1292         time(H_state), *coordinate(H_state).list(), *momentum(H_state).list()
1293     )

```

---

The first test is to see whether  $\{Q, H\} = \partial_2 H$  and  $\{P, H\} = -\partial_1 H$ , where  $Q$  and  $P$  are the coordinate and momentum selectors, and  $H$  is a general state function.

```

1294 q = column_matrix([var("q_x"), var("q_y")])
1295 p = row_matrix([var("p_x"), var("p_y")])
1296 sigma = up(t, q, p)
1297 H = state_function("H")
1298
1299 show(Poisson_bracket(coordinate, H)(sigma))
1300 show(Poisson_bracket(momentum, H)(sigma))

```

---

$$\begin{bmatrix} \frac{\partial}{\partial p_x} H(t, q_x, q_y, p_x, p_y) \\ \frac{\partial}{\partial p_y} H(t, q_x, q_y, p_x, p_y) \end{bmatrix}$$

$$\begin{bmatrix} -\frac{\partial}{\partial q_x} H(t, q_x, q_y, p_x, p_y) \\ -\frac{\partial}{\partial q_y} H(t, q_x, q_y, p_x, p_y) \end{bmatrix}$$

All is correct. Note that both results are standing vectors.

### 3.2.4 Properties of the Poisson bracket

We know that  $\{H, H\} = 0$  for any function. Let's test this for our implementation.

```

1301 show(Poisson_bracket(H, H)(sigma))

```

---

$$[ 0 ]$$

The property  $\{F, F\} = 0$  is actually implied when we can show that the Poisson bracket is anti-symmetric.

---

```

1302 F = state_function("F")
1303 G = state_function("G")
1304
1305 show((Poisson_bracket(F, G) + Poisson_bracket(G, F))(sigma))

```

---

$$[ 0 ]$$

How about  $\{F, G + H\} = \{F, G\} + \{F, H\}$ ?

---

```

1306 show(
1307     (
1308         Poisson_bracket(F, G + H)
1309         - Poisson_bracket(F, G)
1310         - Poisson_bracket(F, H)
1311     )(sigma)
1312 )

```

---

$$[ 0 ]$$

To check the rule  $\{F, cG\} = c\{F, G\}$  we need a constant function. By making the next function independent of any argument, it becomes constant.

---

```

1313 constant = Function(lambda H_state: function("c")())

```

---

Is it indeed constant?

---

```

1314 show(Jacobian(constant)(sigma, sigma))

```

---

$$[ 0 \ 0 \ 0 \ 0 \ 0 ]$$

So, next we can check  $\{F, cG\} = c\{F, G\}$ .

---

```

1315 show(
1316     (Poisson_bracket(F, constant * G) - constant * Poisson_bracket(F, G))(
1317         sigma
1318     ).simplify_full()
1319 )

```

---

$$[ 0 ]$$

Finally, here is the check on Jacobi's identity.

---

```

1320      _____ ./.sage/section3.2.sage _____
1321  jacobi = (
1322      Poisson_bracket(F, Poisson_bracket(G, H))
1323      + Poisson_bracket(G, Poisson_bracket(H, F))
1324      + Poisson_bracket(H, Poisson_bracket(F, G)))
1325  )
1326  show(jacobi(sigma).simplify_full())

```

---

[ 0 ]

### 3.2.5 Poisson bracket of a conserved quantity

To check that the Poisson bracket of a conserved quantity is conserved we need a function that does not depend on time.

---

```

1327  _____ ./.sage/section3.2.sage _____
1328  def f(H_state):
1329      return function("f")(
1330          *coordinate(H_state).list(), *momentum(H_state).list()
1331      )

```

---

Clearly, the derivative with respect to time of this function is zero, so it does what we need.

---

```

1331  _____ ./.sage/section3.2.sage _____
1332  show(diff(f(sigma), time(sigma)))

```

---

0

Now consider  $\{F, H\}$  where  $H$  is the rectangular Hamiltonian.

---

```

1332  _____ ./.sage/section3.2.sage _____
1333  V = Function(lambda q: function("V")(*q.list()))
1334  var(m, domain="positive")
1335
1336  H = H_rectangular(m, V)

```

---

I compute the Poisson bracket of  $F$  and  $H$  for one dimension so that the result remains small.

---

```

1337  _____ ./.sage/section3.2.sage _____
1338  q = column_matrix([var("q")])
1339  p = row_matrix([var("p")])
1340  sigma = up(t, q, p)
1341  show(Poisson_bracket(f, H)(sigma).expand())

```

---

$$\left[ -\frac{\partial}{\partial q} V(q) \frac{\partial}{\partial p} f(q, p) + \frac{p \frac{\partial}{\partial q} f(q, p)}{m} \right]$$

To complete the check, note that, by Hamilton's equation,  $\dot{q} = \partial H / \partial p$ ,  $\dot{p} = -\partial H / \partial q = -\partial V / \partial q$ . If we replace that in the above equation we obtain

$$\dot{p} \frac{\partial f}{\partial p} + \dot{q} \frac{\partial f}{\partial q} = \frac{df}{dt}.$$

Since  $f$  is conserved, the total time derivative of  $F$  is zero, hence  $f$  and  $H$  commute.

## 3.4 PHASE SPACE REDUCTION

### 3.4.1 Standard imports

---

```
1342 load("utils3.1.sage")                                .../sage/section3.2.sage
1343 t = var("t", domain="real")
1344
1345 load("show_expression.sage")                         don't tangle
```

---

### 3.4.2 Motion in a central potential

---

```
1346 var("m")
1347
1348 V = function("V")
1349
1350
1351 def L_polar(m, V):
1352     def Lagrangian(local):
1353         r, phi = coordinate(local).list()
1354         rdot, phidot = velocity(local).list()
1355         T = 1 / 2 * m * (square(rdot) + square(r * phidot))
1356         return T - V(r)
1357
1358     return Lagrangian
```

---



---

```
1359 q = column_path([literal_function("r"), literal_function("phi")])
1360 p = row_path([literal_function("p_r"), literal_function("p_phi")])
1361 H_state = qp_to_H_state_path(q, p)(t)
1362 show(H_state)
```

---

$$\begin{bmatrix} t \\ r \\ \phi \\ [p_r \ p_\phi] \end{bmatrix}$$

---

```
1363 H = Lagrangian_to_Hamiltonian(L_polar(m, V))
1364 show(H(H_state).simplify_full())
```

---

$$\left[ \frac{2mV(r)r^2 + p_r^2 r^2 + p_\phi^2}{2mr^2} \right]$$

Here are the Hamilton equations.

---

```
1365 HE = Hamilton_equations(Lagrangian_to_Hamiltonian(L_polar(m, V)))(q, p)(t)
1366 show(HE)
```

---

$$\begin{bmatrix} 0 \\ -\frac{p_r}{m} + \dot{r} \\ -\frac{p_\phi}{mr^2} + \dot{\phi} \\ -\frac{p_\phi^2}{mr^3} + D_0(V)(r) + \dot{p}_r \quad \dot{p}_\phi \end{bmatrix}$$

Realize, we have obtained the LHS of the system of differential equations  $Dz(t) - F(t, z(t)) = 0$ .

## 3.5 PHASE SPACE EVOLUTION

### 3.5.1 The standard imports.

---

```
.../sage/section3.5.sage
1367 import numpy as np
1368
1369 load("utils1.7.sage", "utils3.1.sage")
1370
1371 var("t", domain="real")
```

---

```
don't tangle
1372 load("show_expression.sage")
```

---

### 3.5.2 The Hamiltonian for the driven pendulum

---

```
.../sage/section3.5.sage
1373 q = column_path([literal_function("theta")])
1374 p = row_path([literal_function(r"p_theta")])
1375 H_state = qp_to_H_state_path(q, p)(t)
1376 show(H_state)
```

---

$$\begin{bmatrix} t \\ \theta \\ p_\theta \end{bmatrix}$$

This is the Hamiltonian. The computations return a  $(1 \times 1)$  matrix; we therefore unpack it.

---

```
..... ./sage/section3.5.sage
1377 _ = var("A g l m omega", domain="positive")
1378
1379 H = Lagrangian_to_Hamiltonian(
1380     L_periodically_driven_pendulum(m, l, g, A, omega)
1381 )
1382 show(H(H_state)[0,0].simplify_full().expand())
```

---

$$-\frac{1}{2} A^2 m \omega^2 \cos(\theta)^2 \sin(\omega t)^2 + A g m \cos(\omega t) - g l m \cos(\theta) + \frac{A \omega p_\theta \sin(\omega t) \sin(\theta)}{l} + \frac{p_\theta^2}{2 l^2 m}$$

Next is the system derivative, i.e., the LHS of the Hamilton equations.

---

```
..... ./sage/section3.5.sage
1383 DH = Hamiltonian_to_state_derivative(H)(H_state)
1384 show(DH[1].simplify_full()[0,0])
1385 show(DH[2].simplify_full()[0,0])
```

---

$$\frac{A l m \omega \sin(\omega t) \sin(\theta) + p_\theta}{l^2 m}$$

$$-\frac{A \omega \cos(\theta) p_\theta \sin(\omega t) + (A^2 l m \omega^2 \cos(\theta) \sin(\omega t)^2 + g l^2 m) \sin(\theta)}{l}$$

The last step is to numerically integrate the HE and make a graph of  $\theta$  and  $p_\theta$ .

---

```
..... ./sage/section3.5.sage
1386 def H_pend_sysder(m, l, g, A, omega):
1387     Hamiltonian = Lagrangian_to_Hamiltonian(
1388         L_periodically_driven_pendulum(m, l, g, A, omega)
1389     )
1390
1391     def f(state):
1392         return Hamiltonian_to_state_derivative(Hamiltonian)(state)
1393
1394     return f
```

---

```
..... ./sage/section3.5.sage
1395 times = strange(0, 100, 0.001, include_endpoint=True)
1396 soln = evolve(
1397     H_pend_sysder(m=1, l=1, g=9.8, A=0.1, omega=2 * sqrt(9.8)),
1398     ics=up(0, column_matrix([1]), row_matrix([0])),
```

---

```

1399     times=times,
1400 )
1401 thetas = principal_value(np.pi)(soln[:, 1])
1402 thetadots = soln[:, 2]
1403 pp = list(zip(thetas, thetadots))
1404 p = points(pp, color='blue', size=3)
1405 p.save(f'../../figures/hamiltonian_driven_pendulum_0.001.png')

```

---

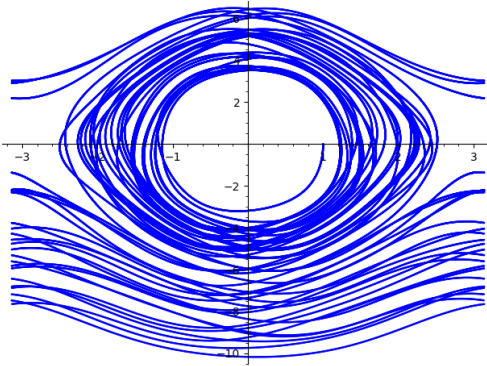


Figure 3.1: The driven pendulum obtained from numerically integrating the Hamilton equations. The graph is not identical to the one in the book, because of the inherent chaotic behavior.

---

```

.../sage/section3.5.sage
1406 times = strange(0, 100, 0.005, include_endpoint=True)
1407 soln = evolve(
1408     H_pend_sysder(m=1, l=1, g=9.8, A=0.1, omega=2 * sqrt(9.8)),
1409     ics=up(0, vector([1]), vector([0])),
1410     times=times,
1411 )
1412 thetas = principal_value(np.pi)(soln[:, 1])
1413 thetadots = soln[:, 2]
1414 pp = list(zip(thetas, thetadots))
1415 p = points(pp, color='blue', size=3)
1416 p.save(f'../../figures/hamiltonian_driven_pendulum_0.005.png')

```

---

### 3.9 THE STANDARD MAP

We take a number of uniformly distributed starting points for the paths. The result, shown in Fig. 3.3, is very nice.

---

```

1417 import numpy as np
1418 import matplotlib.pyplot as plt
1419

```

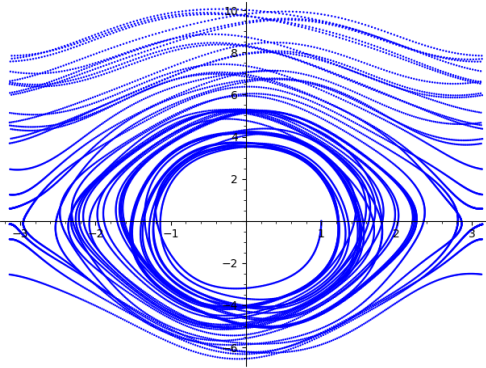


Figure 3.2: The driven pendulum obtained from numerically integrating the Hamilton equations, but now with time steps 0.005 instead of 0.001. The graphs for both step-sizes seem to be qualitatively the same, but the details are different.

```

1420
1421 n_points = 10000
1422 I = np.zeros(n_points)
1423 theta = np.zeros(n_points)
1424
1425 K = 0.9
1426 two_pi = 2 * np.pi
1427
1428 plt.figure(figsize=(10, 10), dpi=300)
1429
1430 for _ in range(500):
1431     theta[0] = np.random.uniform(0, two_pi)
1432     I[0] = np.random.uniform(0, two_pi)
1433     for i in range(1, n_points):
1434         I[i] = (I[i - 1] + K * np.sin(theta[i - 1])) % two_pi
1435         theta[i] = (theta[i - 1] + I[i]) % two_pi
1436         plt.scatter(theta, I, s=0.01, color='black', alpha=0.1, marker='.')
1437
1438
1439 plt.axis('off')
1440 plt.savefig('../figures/standard_map.png', bbox_inches='tight')
```

---

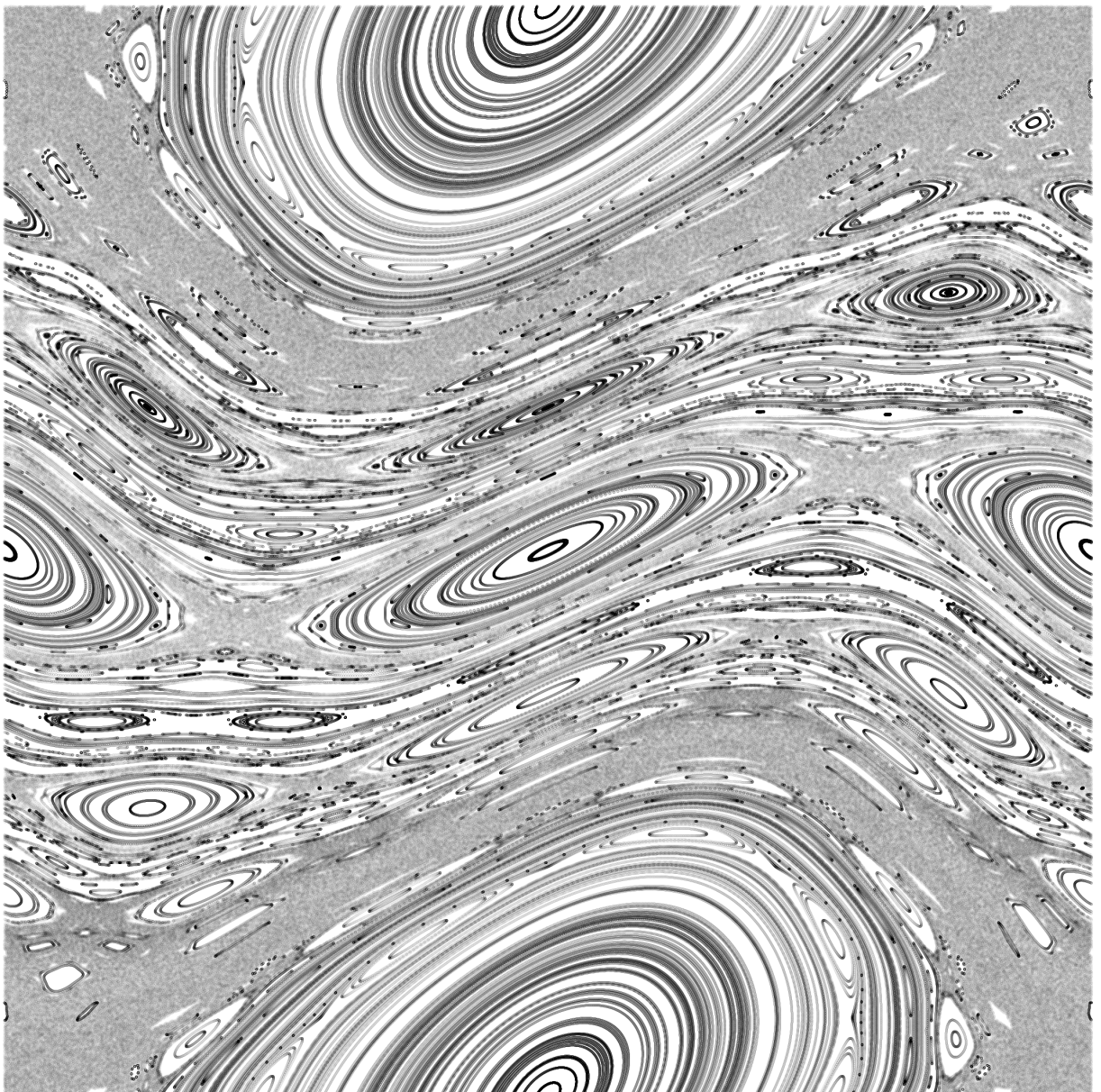


Figure 3.3: The standard map with  $K = 0.6$ .

# 4

CHAPTER 4: SKIPPED

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# 5

## CHAPTER 5

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# 6

## CHAPTER 6

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# 7

## CHAPTER 7

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