distributed Documentation

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Getting Started

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Dask.distributed is a lightweight library for distributed computing in Python. It extends both the concurrent.futures and dask APIs to moderate sized clusters.

See the quickstart to get started.

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2 Getting Started

Motivation

Why build yet-another-distributed-system?

Distributed serves to complement the existing PyData analysis stack. In particular it meets the following needs:

- Low latency: Each task suffers about 1ms of overhead. A small computation and network roundtrip can complete in less than 10ms.
- **Peer-to-peer data sharing:** Workers communicate with each other to share data. This removes central bottle-necks for data transfer.
- **Complex Scheduling:** Supports complex workflows (not just map/filter/reduce) which are necessary for sophisticated algorithms used in nd-arrays, machine learning, image processing, and statistics.
- **Pure Python:** Built in Python using well-known technologies. This eases installation, improves efficiency (for Python users), and simplifies debugging.
- Data Locality: Scheduling algorithms cleverly execute computations where data lives. This minimizes network traffic and improves efficiency.
- Familiar APIs: Compatible with the concurrent futures API in the Python standard library. Compatible with dask API for parallel algorithms
- Easy Setup: As a Pure Python package distributed is pip installable and easy to set up on your own cluster.

Architecture

Dask.distributed is a centrally managed, distributed, dynamic task scheduler. The central dask-scheduler process coordinates the actions of several dask-worker processes spread across multiple machines and the concurrent requests of several clients.

The scheduler is asynchronous and event driven, simultaneously responding to requests for computation from multiple clients and tracking the progress of multiple workers. The event-driven and asynchronous nature makes it flexible to concurrently handle a variety of workloads coming from multiple users at the same time while also handling a fluid worker population with failures and additions. Workers communicate amongst each other for bulk data transfer over sockets.

Internally the scheduler tracks all work as a constantly changing directed acyclic graph of tasks. A task is a Python function operating on Python objects, which can be the results of other tasks. This graph of tasks grows as users submit more computations, fills out as workers complete tasks, and shrinks as users leave or become disinterested in previous results.

Users interact by connecting a local Python session to the scheduler and submitting work, either by individual calls to the simple interface e.submit (function, *args, **kwargs) or by using the large data collections and parallel algorithms of the parent dask library. The collections in the dask library like dask.array and dask.dataframe provide easy access to sophisticated algorithms and familiar APIs like NumPy and Pandas, while the simple e.submit interface provides users with custom control when they want to break out of canned "big data" abstractions and submit fully custom workloads.

Contents

3.1 Install Distributed

You can install distributed with conda, with pip, or by installing from source.

3.1.1 Conda

To install the latest version of distributed from the conda-forge repository using conda:

```
conda install distributed -c conda-forge
```

3.1.2 Pip

Or install distributed with pip:

```
pip install distributed --upgrade
```

3.1.3 Source

To install distributed from source, clone the repository from github:

```
git clone https://github.com/dask/distributed.git
cd distributed
python setup.py install
```

3.1.4 Notes

Note for Macports users: There is a known issue. with python from macports that makes executables be placed in a location that is not available by default. A simple solution is to extend the *PATH* environment variable to the location where python from macports install the binaries:

```
$ export PATH=/opt/local/Library/Frameworks/Python.framework/Versions/3.5/bin:$PATH
or
$ export PATH=/opt/local/Library/Frameworks/Python.framework/Versions/2.7/bin:$PATH
```

3.2 Quickstart

3.2.1 Install

```
$ pip install distributed --upgrade
```

See installation document for more information.

3.2.2 Setup Dask.distributed the Hard Way

Set up scheduler and worker processes on your local computer:

```
$ dask-scheduler
Scheduler started at 127.0.0.1:8786
$ dask-worker 127.0.0.1:8786
$ dask-worker 127.0.0.1:8786
$ dask-worker 127.0.0.1:8786
```

Launch an Executor and point it to the IP/port of the scheduler.

```
>>> from distributed import Executor
>>> executor = Executor('127.0.0.1:8786')
```

See setup for advanced use.

3.2.3 Setup Dask.distributed the Easy Way

If you create an executor without providing an address it will start up a local scheduler and worker for you.

```
>>> from distributed import Executor
>>> executor = Executor()
>>> executor
<Executor: scheduler="127.0.0.1:8786" processes=8 cores=8>
```

Map and Submit Functions

Use the map and submit methods to launch computations on the cluster. The map/submit functions send the function and arguments to the remote workers for processing. They return Future objects that refer to remote data on the cluster. The Future returns immediately while the computations run remotely in the background.

```
>>> def square(x):
    return x ** 2

>>> def neg(x):
    return -x

>>> A = executor.map(square, range(10))
>>> B = executor.map(neg, A)
>>> total = executor.submit(sum, B)
>>> total.result()
-285
```

Gather

The map/submit functions return Future objects, lightweight tokens that refer to results on the cluster. By default the results of computations *stay on the cluster*.

```
>>> total # Function hasn't yet completed
<Future: status: waiting, key: sum-58999c52e0fa35c7d7346c098f5085c7>
>>> total # Function completed, result ready on remote worker
<Future: status: finished, key: sum-58999c52e0fa35c7d7346c098f5085c7>
```

Gather results to your local machine either with the Future.result method for a single future, or with the Executor.gather method for many futures at once.

```
>>> total.result()  # result for single future

-285

>>> executor.gather(A) # gather for many futures

[0, 1, 4, 9, 16, 25, 36, 49, 64, 81]
```

Restart

When things go wrong, or when you want to reset the cluster state, call the restart method.

```
>>> executor.restart()
```

See executor for advanced use.

3.3 Setup Network

A distributed network consists of one Scheduler node and several Worker nodes. One can set these up in a variety of ways

3.3.1 Using the Command Line

We launch the dask-scheduler executable in one process and the dask-worker executable in several processes, possibly on different machines.

Launch dask-scheduler on one node:

```
$ dask-scheduler
Start scheduler at 192.168.0.1:8786
```

Then launch dask-worker on the rest of the nodes, providing the address to the node that hosts dask-scheduler:

```
$ dask-worker 192.168.0.1:8786

Start worker at: 192.168.0.2:12345

Registered with center at: 192.168.0.1:8786

$ dask-worker 192.168.0.1:8786

Start worker at: 192.168.0.3:12346

Registered with center at: 192.168.0.1:8786

$ dask-worker 192.168.0.1:8786
```

```
Start worker at: 192.168.0.4:12347
Registered with center at: 192.168.0.1:8786
```

There are various mechanisms to deploy these executables on a cluster, ranging from manualy SSH-ing into all of the nodes to more automated systems like SGE/SLURM/Torque or Yarn/Mesos.

3.3.2 Using SSH

For this functionality, paramiko library must be installed (e.g. by running pip install paramiko).

The convenience script dask-ssh opens several SSH connections to your target computers and initializes the network accordingly. You can give it a list of hostnames or IP addresses:

```
$ dask-ssh 192.168.0.1 192.168.0.2 192.168.0.3 192.168.0.4
```

Or you can use normal UNIX grouping:

```
$ dask-ssh 192.168.0.{1,2,3,4}
```

Or you can specify a hostfile that includes a list of hosts:

```
$ cat hostfile.txt
192.168.0.1
192.168.0.2
192.168.0.3
192.168.0.4
$ dask-ssh --hostfile hostfile.txt
```

3.3.3 Using the Python API

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Alternatively you can start up the distributed.scheduler.Scheduler and distributed.worker.Worker objects within a Python session manually. Both are torando.tcpserver.TCPServer objects.

Start the Scheduler, provide the listening port (defaults to 8786) and Tornado IOLoop (defaults to IOLoop.current())

```
from distributed import Scheduler
s = Scheduler(loop=loop)
s.start(port)
```

On other nodes start worker processes that point to the IP address and port of the scheduler.

```
from distributed import Worker
w = Worker('192.168.0.1', 8786, loop=loop)
w.start(0) # choose randomly assigned port
```

Alternatively, replace Worker with Nanny if you want your workers to be managed in a separate process by a local nanny process.

If you do not already have a Tornado event loop running you will need to create and start one, possibly in a separate thread.

```
from tornado.ioloop import IOLoop
loop = IOLoop()
```

```
from threading import Thread
t = Thread(target=loop.start)
t.start()
```

3.3.4 Using Amazon EC2

See the EC2 quickstart for information on the dask-ec2 easy setup script to launch a canned cluster on EC2.

3.3.5 Cleanup

It is common and safe to terminate the cluster by just killing the processes. The workers and scheduler have no persistent state.

Programmatically you can use the client interface (rpc) to call the terminate methods on the workers and schedulers.

3.4 EC2 Startup Script

First, add your AWS credentials to ~/.aws/credentials like this:

```
[default]
aws_access_key_id = YOUR_ACCESS_KEY
aws_secret_access_key = YOUR_SECRET_KEY
```

For other ways to manage or troubleshoot credentials, see the boto3 docs.

Now, you can quickly deploy a scheduler and workers on EC2 using the dask-ec2 quickstart application:

```
pip install dask-ec2
dask-ec2 up --keyname YOUR-AWS-KEY --keypair ~/.ssh/YOUR-AWS-SSH-KEY.pem
```

This provisions a cluster on Amazon's EC2 cloud service, installs Anaconda, and sets up a scheduler and workers. In then prints out instructions on how to connect to the cluster.

3.4.1 Options

The dask-ec2 startup script comes with the following options for creating a cluster:

```
$ dask-ec2 up --help
Usage: dask-ec2 up [OPTIONS]
Options:
 --keyname TEXT
                               Keyname on EC2 console [required]
 --keypair PATH
                               Path to the keypair that matches the keyname [required]
 --name TEXT
                               Tag name on EC2
  --region-name TEXT
                               AWS region [default: us-east-1]
  --ami TEXT
                               EC2 AMI [default: ami-d05e75b8]
 --username TEXT
                               User to SSH to the AMI [default: ubuntu]
 --type TEXT
                               EC2 Instance Type [default: m3.2xlarge]
                               Number of nodes [default: 4]
 --count INTEGER
                               Security Group Name [default: dask-ec2-default]
 --security-group TEXT
 --volume-type TEXT
                               Root volume type [default: gp2]
 --volume-size INTEGER
                               Root volume size (GB) [default: 500]
```

```
--file PATH File to save the metadata [default: cluster.yaml]
--provision / --no-provision Provision salt on the nodes [default: True]
--dask / --no-dask Install Dask.Distributed in the cluster [default: True]
--nprocs INTEGER Number of processes per worker [default: 1]
-h, --help Show this message and exit.
```

3.4.2 Connect

Connection instructions follow successful completion of the dask-ec2 up command. The involve the following:

```
dask-ec2 ssh  # SSH into head node
ipython  # Start IPython console on head node
```

```
>>> from distributed import Executor, s3, progress
>>> e = Executor('127.0.0.1:8786')
```

This executor now has access to all the cores of your cluster.

3.4.3 Destroy

You can destroy your cluster from your local machine with the destroy command:

```
dask-ec2 destroy
```

3.5 Web Interface

Information about the current state of the network helps to track progress, identify performance issues, and debug failures.

Dask.distributed includes a web interface to help deliver this information over a normal web page in real time. This web interface is launched by default wherever the scheduler is launched if the scheduler machine has Bokeh installed (conda install bokeh). The web interface is normally available at http://scheduler-address:8787/status/ and can be viewed any normal web browser.

The web UI shows basic statistics on all worker machines, grouped by physical address. This includes information like CPU/memory load, active tasks, latency, network and disk usage, etc.. The tabular statistics are updated about once a second.

It also shows the progress of all groups of tasks currently running on the cluster. Dark blue is used for tasks that are completed and in memory, light blue for tasks that are completed and have been released, gray for not yet completed, and black for erred. The progress bar is updated every 100ms.

There is a resource plot showing total CPU and memory use of the cluster over the last few minutes.

Finally there is a plot of tasks as they complete, showing their start and end times, start and end transfer times (in red), as well as which worker they were run on. Hovering over any of the tasks gives the task name as well as more precise information. This plot can be invaluable to determine performance issues. It is updated every 200ms. It only includes the most recent thousand tasks. For the most recent 20000 tasks visit http://my-scheduler-address:8787/tasks , although beware that this page is not updated in real time.

3.6 Examples

3.6.1 Word count in HDFS

Setup

In this example, we'll use distributed with the hdfs3 library to count the number of words in text files (Enron email dataset, 6.4 GB) stored in HDFS.

Copy the text data from Amazon S3 into HDFS on the cluster:

```
$ hadoop distcp s3n://AWS_SECRET_ID:AWS_SECRET_KEY@blaze-data/enron-email hdfs:///tmp/enron
```

where AWS_SECRET_ID and AWS_SECRET_KEY are valid AWS credentials.

Start the distributed scheduler and workers on the cluster.

Code example

Import distributed, hdfs3, and other standard libraries used in this example:

```
>>> import hdfs3
>>> from collections import defaultdict, Counter
>>> from distributed import Executor, progress
```

Initalize a connection to HDFS, replacing NAMENODE_HOSTNAME and NAMENODE_PORT with the hostname and port (default: 8020) of the HDFS namenode.

```
>>> hdfs = hdfs3.HDFileSystem('NAMENODE_HOSTNAME', port=NAMENODE_PORT)
```

Initalize a connection to the distributed executor, replacing EXECUTOR_IP and EXECUTOR_PORT with the IP address and port of the distributed scheduler.

```
>>> e = Executor('EXECUTOR_IP:EXECUTOR_PORT')
```

Generate a list of filenames from the text data in HDFS:

Print the first 1024 bytes of the first text file:

```
b'Date: Wed, 29 Nov 2000 09:33:00 -0800 (PST)\r\nFrom: Xochitl-Alexis Velasc
o\r\nTo: Mark Knippa, Mike D Smith, Gerald Nemec, Dave S Laipple, Bo Barnwel
l\r\nCc: Melissa Jones, Iris Waser, Pat Radford, Bonnie Shumaker\r\nSubject:
Finalize ECS/EES Master Agreement\r\nX-SDOC: 161476\r\nX-ZLID: zl-edrm-enro
n-v2-nemec-g-2802.eml\r\n\r\nPlease plan to attend a meeting to finalize the
ECS/EES Master Agreement \r\ntomorrow 11/30/00 at 1:30 pm CST.\r\n\r\nI wi
ll email everyone tomorrow with location.\r\n\r\nDave-I will also email you
the call in number tomorrow.\r\n\r\nThanks\r\nXochitl\r\n\r\n***********
```

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```
EDRM Enron Email Data Set has been produced in EML, PST and NSF format by ZL Technologies, Inc. This Data Set is licensed under a Creative Commons Attri bution 3.0 United States License <a href="http://creativecommons.org/licenses/by/3.0/us/">http://creativecommons.org/licenses/by/3.0/us/</a>. To provide attribution, please cite to "ZL Technologies, Inc. (http://www.zlti.com)."\r\n********\r\nDate: Wed, 29 Nov 2000 09:40:00 -0800 (PST)\r\nFrom: Jill T Zivley\r\nTo: Robert Cook, Robert Crockett, John Handley, Shawna'
```

Create a function to count words in each file:

Before we process all of the text files using the distributed workers, let's test our function locally by counting the number of words in the first text file:

```
>>> counts = count_words(filenames[0])
>>> print(sorted(counts.items(), key=lambda k_v: k_v[1], reverse=True)[:10])

[(b'the', 144873),
(b'of', 98122),
(b'to', 97202),
(b'and', 90575),
(b'or', 60305),
(b'in', 53869),
(b'a', 43300),
(b'any', 31632),
(b'by', 31515),
(b'is', 30055)]
```

We can perform the same operation of counting the words in the first text file, except we will use e.submit to execute the computation on a distributed worker:

```
>>> future = e.submit(count_words, filenames[0])
>>> counts = future.result()
>>> print(sorted(counts.items(), key=lambda k_v: k_v[1], reverse=True)[:10])

[(b'the', 144873),
   (b'of', 98122),
   (b'to', 97202),
   (b'and', 90575),
   (b'or', 60305),
   (b'in', 53869),
   (b'a', 43300),
   (b'any', 31632),
   (b'by', 31515),
   (b'is', 30055)]
```

We are ready to count the number of words in all of the text files using distributed workers. Note that the map operation is non-blocking, and you can continue to work in the Python shell/notebook while the computations are running.

```
>>> futures = e.map(count_words, filenames)
```

We can check the status of some futures while all of the text files are being processed:

When the futures finish reading in all of the text files and counting words, the results will exist on each worker. This operation required about 3 minutes to run on a cluster with three worker machines, each with 4 cores and 16 GB RAM.

Note that because the previous computation is bound by the GIL in Python, we can speed it up by starting the distributed workers with the --nprocs 4 option.

To sum the word counts for all of the text files, we need to gather some information from the distributed workers. To reduce the amount of data that we gather from the workers, we can define a function that only returns the top 10,000 words from each text file.

```
>>> def top_items(d):
...    items = sorted(d.items(), key=lambda kv: kv[1], reverse=True)[:10000]
...    return dict(items)
```

We can then map the futures from the previous step to this culling function. This is a convenient way to construct a pipeline of computations using futures:

```
>>> futures2 = e.map(top_items, futures)
```

We can gather the resulting culled word count data for each text file to the local process:

```
>>> results = e.gather(iter(futures2))
```

To sum the word counts for all of the text files, we can iterate over the results in futures2 and update a local dictionary that contains all of the word counts.

```
>>> all_counts = Counter()
>>> for result in results:
... all_counts.update(result)
```

Finally, we print the total number of words in the results and the words with the highest frequency from all of the text files:

```
>>> print(len(all_counts))

8797842

>>> print(sorted(all_counts.items(), key=lambda k_v: k_v[1], reverse=True)[:10])

[(b'0', 67218380),
   (b'the', 19586868),
   (b'-', 14123768),
   (b'to', 11893464),
```

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```
(b'N/A', 11814665),
(b'of', 11724827),
(b'and', 10253753),
(b'in', 6684937),
(b'a', 5470371),
(b'or', 5227805)]
```

The complete Python script for this example is shown below:

```
# word-count.py
import hdfs3
from collections import defaultdict, Counter
from distributed import Executor, progress
hdfs = hdfs3.HDFileSystem('NAMENODE_HOSTNAME', port=NAMENODE_PORT)
e = Executor('EXECUTOR_IP:EXECUTOR_PORT')
filenames = hdfs.glob('/tmp/enron/*/*')
print(filenames[:5])
print (hdfs.head(filenames[0]))
def count_words(fn):
   word_counts = defaultdict(int)
   with hdfs.open(fn) as f:
        for line in f:
            for word in line.split():
                word_counts[word] += 1
   return word_counts
counts = count_words(filenames[0])
print(sorted(counts.items(), key=lambda k_v: k_v[1], reverse=True)[:10])
future = e.submit(count_words, filenames[0])
counts = future.result()
print(sorted(counts.items(), key=lambda k_v: k_v[1], reverse=True)[:10])
futures = e.map(count_words, filenames)
len (futures)
futures[:5]
progress (futures)
def top_items(d):
   items = sorted(d.items(), key=lambda kv: kv[1], reverse=True)[:10000]
   return dict(items)
futures2 = e.map(top_items, futures)
results = e.gather(iter(futures2))
all_counts = Counter()
for result in results:
    all_counts.update(result)
print (len(all_counts))
print(sorted(all_counts.items(), key=lambda k_v: k_v[1], reverse=True)[:10])
```

3.7 Executor

The Executor is the primary entry point for users of distributed.

After you setup a cluster, initialize an Executor by pointing it to the address of a Scheduler:

```
>>> from distributed import Executor
>>> executor = Executor('127.0.0.1:8786')
```

3.7.1 **Usage**

submit

You can submit individual function calls with the executor.submit method

The result is on one of the distributed workers. We can continue using x in further calls to submit:

```
>>> type(x)
Future
>>> y = executor.submit(inc, x)
```

Gather results

We can collect results in a variety of ways. First, we can use the .result () method on futures

```
>>> x.result()
2
```

Second, we can use the gather method on the executor

```
>>> executor.gather([x, y])
(2, 3)
```

Third, we can use the as_completed function to iterate over results as soon as they become available.

```
>>> from distributed import as_completed
>>> seq = as_completed([x, y])
>>> next(seq).result()
2
>>> next(seq).result()
3
```

But, as always, we want to minimize communicating results back to the local process. It's often best to leave data on the cluster and operate on it remotely with functions like submit, map, get and compute. See efficiency for more information on efficient use of distributed.

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map

We can map a function over many inputs at once

```
>>> L = executor.map(inc, range(10))
```

The map method returns a list of futures. This is a break with the concurrent futures API, which returns the results directly. We keep the results as futures so that they can stay on the distributed cluster.

Additionally, we don't do any kind of batching so every function application will be a new task which will have a couple milliseconds of overhead. It is unwise to use executor.map for small, fast functions where scheduling overhead is likely to be more expensive than the cost of the function itself. For example, our function inc is actually a *terrible* function to parallelize in practice.

dask

Distributed provides a dask compliant task scheduling interface. It provides this through two methods, get (synchronous) and compute (asynchronous).

get

We provide dask graph dictionaries to the scheduler:

```
>>> dsk = {'x': 1, 'y': (inc, 'x')}
>>> executor.get(dsk, 'y')
2
```

This function pulls results back by default. This is so that it can integrate with existing dask code.

```
>>> import dask.array as da

>>> x = da.random.random(1000000000, chunks=(1000000,))

>>> x.sum().compute() # use local threads

499999359.23511785

>>> x.sum().compute(get=executor.get) # use distributed cluster

499999359.23511785
```

compute

We can also provide dask collections (arrays, bags, dataframes, delayed values) to the executor with the compute method.

```
>>> type(x)
dask.array.Array
>>> type(df)
dask.dataframe.DataFrame
>>> x_future, df_future = executor.compute(x, df)
```

This immediately returns standard Future objects as would be returned by submit or map.

restart

When things go wrong, restart the cluster with the .restart () method.

```
>>> executor.restart()
```

This both resets the scheduler state and all of the worker processes. All current data and computations will be lost. All existing futures set their status to 'cancelled'.

See resilience for more information.

3.7.2 Internals

Data Locality

By default the executor does not bring results back to your local computer but leaves them on the distributed network. As a result, computations on returned results like the following don't require any data transfer.

```
>>> y = executor.submit(inc, x) # no data transfer required
```

In addition, the internal scheduler endeavors to run functions on worker nodes that already have the necessary input data. It avoids worker-to-worker communication when convenient.

Pure Functions by Default

By default we assume that all functions are pure. If this is not the case you should use the pure=False keyword argument.

The executor associates a key to all computations. This key is accessible on the Future object.

```
>>> from operator import add

>>> x = executor.submit(add, 1, 2)

>>> x.key

'add-ebf39f96ad7174656f97097d658f3fa2'
```

This key should be the same accross all computations with the same inputs and across all machines. If you run the computation above on any computer with the same environment then you should get the exact same key.

The scheduler avoids redundant computations. If the result is already in memory from a previous call then that old result will be used rather than recomputing it. Calls to submit or map are idempotent in the common case.

While convenient, this feature may be undesired for impure functions, like random. In these cases two calls to the same function with the same inputs should produce different results. We accomplish this with the pure=False keyword argument. In this case keys are randomly generated (by uuid4.)

```
>>> import numpy as np
>>> executor.submit(np.random.random, 1000, pure=False).key
'random_sample-fc814a39-ee00-42f3-8b6f-cac65bcb5556'
>>> executor.submit(np.random.random, 1000, pure=False).key
'random_sample-a24e7220-a113-47f2-a030-72209439f093'
```

Garbage Collection

Prolonged use of distributed may allocate a lot of remote data. The executor can clean up unused results by reference counting.

The executor reference counts Future objects. When a particular key no longer has any Future objects pointing to it it will be released from distributed memory if no active computations still require it.

In this way garbage collection in the distributed memory space of your cluster mirrors garbage collection within your local Python session.

Known future keys and reference counts can be found in the following dictionaries:

```
>>> executor.futures
>>> executor.refcount
```

The scheduler also cleans up intermediate results when provided full dask graphs. You can always use the lower level delete or clear functions in distributed.client to manage data manually.

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Dask Graph

The executor and scheduler maintain a dask graph of all known computations. This graph is accessible via the .dask attribute. At times it may be worth visualizing this object.

```
>>> executor.dask
>>> from dask.base import visualize
>>> visualize(executor, filename='executor.pdf')
```

All functions like .submit, .map, and .get just add small subgraphs to this graph. Functions like .result, as_completed, or .gather, wait until their respective parts of the graph have completed. All of these actions are asynchronous to the actual execution of the graph, which is managed in a background thread.

The dask graph is also used to recover results in case of failure.

Coroutines

If you are operating in an asynchronous environment then all blocking functions listed above have asynchronous equivalents. Currently these have the exact same name but are prepended with an underscore (_) so, .result is synchronous while ._result is asynchronous. If a function has no asynchronous counterpart then that means it does not significantly block. The .submit and .map functions are examples of this; they return immediately in either case.

3.8 Local Cluster

For convenience you can start a local cluster from your Python session.

```
>>> from distributed import Executor, LocalCluster
>>> c = LocalCluster()
LocalCluster("127.0.0.1:8786", workers=8, ncores=8)
>>> e = Executor(c)
<Executor: scheduler=127.0.0.1:8786 processes=8 cores=8>
```

Alternatively, a LocalCluster is made for you automatically if you create an Executor with no arguments.

```
>>> from distributed import Executor
>>> e = Executor()
>>> e
<Executor: scheduler=127.0.0.1:8786 processes=8 cores=8>
```

Create local Scheduler and Workers

This creates a "cluster" of a scheduler and workers running on the local machine.

```
Parameters n_workers: int
```

Number of workers to start

threads per worker: int

Number of threads per each worker

nanny: boolean

If true start the workers in separate processes managed by a nanny. If False keep the workers in the main calling process

scheduler_port: int

Port of the scheduler. 8786 by default, use 0 to choose a random port

Examples

```
>>> c = LocalCluster() # Create a local cluster with as many workers as cores
>>> c
LocalCluster("192.168.1.141:8786", workers=8, ncores=8)
```

```
>>> e = Executor(c) # connect to local cluster
```

Add a new worker to the cluster >>> w = c.start_worker(ncores=2) # doctest: +SKIP

Shut down the extra worker >>> c.remove_worker(w) # doctest: +SKIP

Start a diagnostic web server and open a new browser tab >>> c.start_diagnostics_server(show=True) # doctest: +SKIP

close()

Close the cluster

```
start_diagnostics_server (port=8787, show=False, silence=50)
```

Start Diagnostics Web Server

This starts a web application to show diagnostics of what is happening on the cluster. This application runs in a separate process and is generally available at the following location:

http://localhost:8787/status/

```
start_worker (port=0, ncores=0, **kwargs)
```

Add a new worker to the running cluster

Parameters port: int (optional)

Port on which to serve the worker, defaults to 0 or random

```
ncores: int (optional)
```

Number of threads to use. Defaults to number of logical cores

nanny: boolean

If true start worker in separate process managed by a nanny

Returns The created Worker or Nanny object. Can be discarded.

Examples

```
>>> c = LocalCluster()
>>> c.start_worker(ncores=2)
```

```
stop_worker(w)
```

Stop a running worker

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Examples

```
>>> c = LocalCluster()
>>> w = c.start_worker(ncores=2)
>>> c.stop_worker(w)
```

3.9 Efficiency

Parallel computing done well is responsive and rewarding. However, several speed-bumps can get in the way. This section describes common ways to ensure performance.

3.9.1 Leave data on the cluster

Wait as long as possible to gather data locally. If you want to ask a question of a large piece of data on the cluster it is often faster to submit a function onto that data then to bring the data down to your local computer.

For example if we have a numpy array on the cluster and we want to know its shape we might choose one of the following options:

- 1. Slow: Gather the numpy array to the local process, access the .shape attribute
- 2. **Fast:** Send a lambda function up to the cluster to compute the shape

```
>>> x = executor.submit(np.random.random, (1000, 1000))
>>> type(x)
Future
```

Slow

```
>>> x.result().shape() # Slow from lots of data transfer (1000, 1000)
```

Fast

```
>>> executor.submit(lambda a: a.shape, x).result() # fast (1000, 1000)
```

3.9.2 Use larger tasks

The scheduler adds about *one millisecond* of overhead per task or Future object. While this may sound fast it's quite slow if you run a billion tasks. If your functions run faster than 100ms or so then you might not see any speedup from using distributed computing.

A common solution is to batch your input into larger chunks.

Slow

```
>>> futures = executor.map(f, seq)
>>> len(futures) # avoid large numbers of futures
1000000000
```

Fast

```
>>> def f_many(chunk):
...    return [f(x) for x in chunk]

>>> from toolz import partition_all
>>> chunks = partition_all(1000000, seq) # Collect into groups of size 1000

>>> futures = executor.map(f_many, chunks)
>>> len(futures) # Compute on larger pieces of your data at once
1000
```

3.9.3 Adjust between Threads and Processes

By default a single Worker runs many computations in parallel using as many threads as your compute node has cores. When using pure Python functions this may not be optimal and you may instead want to run several separate worker processes on each node, each using one thread. When configuring your cluster you may want to use the options to the dask-worker executable as follows:

```
$ dask-worker ip:port --nprocs 8 --nthreads 1
```

Note that if you're primarily using NumPy, Pandas, SciPy, Scikit Learn, Numba, or other C/Fortran/LLVM/Cython-accelerated libraries then this is not an issue for you. Your code is likely optimal for use with multi-threading.

3.9.4 Don't go distributed

Consider the dask and concurrent.futures modules, which have similar APIs to distributed but operate on a single machine. It may be that your problem performs well enough on a laptop or large workstation.

Consider accelerating your code through other means than parallelism. Better algorithms, data structures, storage formats, or just a little bit of C/Fortran/Numba code might be enough to give you the 10x speed boost that you're looking for. Parallelism and distributed computing are expensive ways to accelerate your application.

3.10 Data Locality

Data movement often needlessly limits performance.

This is especially true for analytic computations. Distributed minimizes data movement when possible and enables the user to take control when necessary. This document describes current scheduling policies and user API around data locality.

3.10.1 Current Policies

Task Submission

In the common case distributed runs tasks on workers that already hold dependent data. If you have a task f(x) that requires some data x then that task will very likely be run on the worker that already holds x.

If a task requires data split among multiple workers, then the scheduler chooses to run the task on the worker that requires the least data transfer to it. The size of each data element is measured by the workers using the sys.getsizeof function, which depends on the __sizeof__ protocol generally available on most relevant Python objects.

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Data Scatter

When a user scatters data from their local process to the distributed network this data is distributed in a round-robin fashion grouping by number of cores. So for example If we have two workers Alice and Bob, each with two cores and we scatter out the list range (10) as follows:

```
futures = e.scatter(range(10))
```

Then Alice and Bob receive the following data

```
• Alice: [0, 1, 4, 5, 8, 9]
```

• Bob: [2, 3, 6, 7]

3.10.2 User Control

Complex algorithms may require more user control.

For example the existence of specialized hardware such as GPUs or database connections may restrict the set of valid workers for a particular task.

In these cases use the workers= keyword argument to the submit, map, or scatter functions, providing a hostname, IP address, or alias as follows:

```
future = e.submit(func, *args, workers=['Alice'])
```

```
• Alice: [0, 1, 4, 5, 8, 9, new_result]
```

• Bob: [2, 3, 6, 7]

Required data will always be moved to these workers, even if the volume of that data is significant. If this restriction is only a preference and not a strict requirement, then add the allow_other_workers keyword argument to signal that in extreme cases such as when no valid worker is present, another may be used.

Additionally the scatter function supports a broadcast= keyword argument to enforce that the all data is sent to all workers rather than round-robined. If new workers arrive they will not automatically receive this data.

```
futures = e.scatter([1, 2, 3], broadcast=True) # send data to all workers
```

```
• Alice: [1, 2, 3]
```

• Bob: [1, 2, 3]

Valid arguments for workers= include the following:

• A single IP addresses, IP/Port pair, or hostname like the following:

```
192.168.1.100, 192.168.1.100:8989, alice, alice:8989
```

• A list or set of the above:

```
['alice'], ['192.168.1.100', '192.168.1.101:9999']
```

If only a hostname or IP is given then any worker on that machine will be considered valid. Additionally, you can provide aliases to workers upon creation.:

```
$ dask-worker scheduler_address:8786 --name worker_1
```

And then use this name when specifying workers instead.

```
e.map(func, sequence, workers='worker_1')
```

See the efficiency page to learn about best practices.

3.11 Memory Management

Dask.distributed stores the results of tasks in the distributed memory of the worker nodes. The central scheduler tracks all data on the cluster and determines when data should be freed. Completed results are usually cleared from memory as quickly as possible in order to make room for more computation. The result of a task is kept in memory if either of the following conditions hold:

- 1. A client holds a future pointing to this task. The data should stay in RAM so that the client can gather the data on demand.
- 2. The task is necessary for ongoing computations that are working to produce the final results pointed to by futures. These tasks will be removed once no ongoing tasks require them.

When users hold Future objects or persisted collections (which contain many such Futures inside their .dask attribute) they pin those results to active memory. When the user deletes futures or collections from their local Python process the scheduler removes the associated data from distributed RAM. Because of this relationship, distributed memory reflects the state of local memory. A user may free distributed memory on the cluster by deleting persisted collections in the local session.

3.11.1 Creating Futures

The following functions produce Futures

Executor.submit(func, *args, **kwargs)	Submit a function application to the scheduler
<pre>Executor.map(func, *iterables, **kwargs)</pre>	Map a function on a sequence of arguments
Executor.compute(args[, sync])	Compute dask collections on cluster
Executor.persist(collections)	Persist dask collections on cluster
<pre>Executor.scatter(data[, workers, broadcast,])</pre>	Scatter data into distributed memory

Submit and map handle raw Python functions. Compute and persist handle Dask collections like arrays, bags, delayed values, and dataframes. Scatter sends data directly from the local process.

3.11.2 Persisting Collections

Calls to Executor.compute or Executor.persist submit task graphs to the cluster and return Future objects that point to particular output tasks.

Compute returns a single future per input, persist returns a copy of the collection with each block or partition replaced by a single future. In short, use persist to keep full collection on the cluster and use compute when you want a small result as a single future.

Persist is more common and is often used as follows with collections:

```
>>> # Construct dataframe, no work happens
>>> df = dd.read_csv(...)
>>> df = df[df.x > 0]
>>> df = df.assign(z = df.x + df.y)
>>> # Pin data in distributed ram, this triggers computation
```

```
>>> df = e.persist(df)
>>> # continue operating on df
```

Note for Spark users: this differs from what you're accustomed to. Persist is an immediate action. However, you'll get control back immediately as computation occurs in the background.

In this example we build a computation by parsing CSV data, filtering rows, and then adding a new column. Up until this point all work is lazy; we've just built up a recipe to perform the work as a graph in the df object.

When we call df = e.perist(df) we cut this graph off of the df object, send it up to the scheduler, receive Future objects in return and create a new dataframe with a very shallow graph that points directly to these futures. This happens more or less immediately (as long as it takes to serialize and send the graph) and we can continue working on our new df object while the cluster works to evaluate the graph in the background.

3.11.3 Difference with dask.compute

The operations e.persist(df) and e.compute(df) are asynchronous and so differ from the traditional df.compute() method or dask.compute function, which blocks until a result is available. The .compute() method does not persist any data on the cluster. The .compute() method also brings the entire result back to the local machine, so it is unwise to use it on large datasets. However, .compute() is very convenient for smaller results particularly because it does return concrete results in a way that most other tools expect.

Typically we use asynchronous methods like e.persist to set up large collections and then use df.compute() for fast analyses.

```
>>> # df.compute() # This is bad and would likely flood local memory
>>> df = e.persist(df) # This is good and asynchronously pins df
>>> df.x.sum().compute() # This is good because the result is small
>>> future = e.compute(df.x.sum()) # This is also good but less intuitive
```

3.11.4 Clearing data

We remove data from distributed ram by removing the collection from our local process. Remote data is removed once all Futures pointing to that data are removed from all client machines.

```
>>> del df # Deleting local data often deletes remote data
```

If this is the only copy then this will likely trigger the cluster to delete the data as well.

However if we have multiple copies or other collections based on this one then we'll have to delete them all.

```
>>> df2 = df[df.x < 10]
>>> del df # would not delete data, because df2 still tracks the futures
```

3.11.5 Aggressively Clearing Data

To definitely remove a computation and all computations that depend on it you can always cancel the futures/collection.

```
>>> e.cancel(df) # kills df, df2, and every other dependent computation
```

Alternatively, if you want a clean slate, you can restart the cluster. This clears all state and does a hard restart of all worker processes. It generally completes in around a second.

```
>>> e.restart()
```

3.11.6 Resilience

Results are not intentionally copied unless necessary for computations on other worker nodes. Resilience is achieved through recomputation by maintaining the provenance of any result. If a worker node goes down the scheduler is able to recompute all of its results. The complete graph for any desired Future is maintained until no references to that future exist.

3.11.7 Advanced techniques

At first the result of a task is not intentionally copied, but only persists on the node where it was originally computed or scattered. However result may be copied to another worker node in the course of normal computation if that result is required by another task that is intended to by run by a different worker. This occurs if a task requires two pieces of data on different machines (at least one must move) or through work stealing. In these cases it is the policy for the second machine to maintain its redundant copy of the data. This helps to organically spread around data that is in high demand.

However, advanced users may want to control the location, replication, and balancing of data more directly throughout the cluster. They may know ahead of time that certain data should be broadcast throughout the network or that their data has become particularly imbalanced, or that they want certain pieces of data to live on certain parts of their network. These considerations are not usually necessary.

Executor.rebalance([futures, workers])	Rebalance data within network	
<pre>Executor.replicate(futures[, n, workers,])</pre>	Set replication of futures within network	
<pre>Executor.scatter(data[, workers, broadcast,])</pre>	Scatter data into distributed memory	

3.12 Joblib Frontend

Joblib is a library for simple parallel programming primarily developed and used by the Scikit Learn community. As of version 0.10 it contains a plugin mechanism to allow Joblib code to use other parallel frameworks to execute computations. The distributed scheduler implements such a plugin in the distributed.joblib module and registers it appropriately with Joblib. As a result, any joblib code (including many scikit-learn algorithms) will run on the distributed scheduler if you enclose it in a context manager as follows:

```
import distributed.joblib
from joblib import Parallel, parallel_backend
with parallel_backend('distributed', scheduler_host='HOST:PORT'):
    # normal Joblib code
```

For example you might distributed a randomized cross validated parameter search as follows:

```
import distributed.joblib
from joblib import Parallel, parallel_backend
from sklearn.datasets import load_digits
from sklearn.grid_search import RandomizedSearchCV
from sklearn.svm import SVC
import numpy as np

digits = load_digits()
```

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```
param_space = {
    'C': np.logspace(-6, 6, 13),
    'gamma': np.logspace(-8, 8, 17),
    'tol': np.logspace(-4, -1, 4),
    'class_weight': [None, 'balanced'],
}

model = SVC(kernel='rbf')
search = RandomizedSearchCV(model, param_space, cv=3, n_iter=50, verbose=10)

with parallel_backend('distributed', scheduler_host='localhost:8786'):
    search.fit(digits.data, digits.target)
```

3.13 Data Streams with Queues

The Executor methods scatter, map, and gather can consume and produce standard Python Queue objects. This is useful for processing continuous streams of data. However, it does not constitute a full streaming data processing pipeline like Storm.

3.13.1 Example

We connect to a local Executor.

```
>>> from distributed import Executor
>>> e = Executor('127.0.0.1:8786')
>>> e
<Executor: scheduler=127.0.0.1:8786 workers=1 threads=4>
```

We build a couple of toy data processing functions:

```
from time import sleep
from random import random

def inc(x):
    from random import random
    sleep(random() * 2)
    return x + 1

def double(x):
    from random import random
    sleep(random())
    return 2 * x
```

And we set up an input Queue and map our functions across it.

```
>>> from queue import Queue
>>> input_q = Queue()
>>> remote_q = e.scatter(input_q)
>>> inc_q = e.map(inc, remote_q)
>>> double_q = e.map(double, inc_q)
```

We will fill the input_q with local data from some stream, and then remote_q, inc_q and double_q will fill with Future objects as data gets moved around.

We gather the futures from the double_q back to a queue holding local data in the local process.

```
>>> result_q = e.gather(double_q)
```

Insert Data Manually

Because we haven't placed any data into any of the queues everything is empty, including the final output, result_q.

```
>>> result_q.qsize()
0
```

But when we insert an entry into the input_q, it starts to make its way through the pipeline and ends up in the result_q.

```
>>> input_q.put(10)
>>> result_q.get()
22
```

Insert data in a separate thread

We simulate a slightly more realistic situation by dumping data into the input_q in a separate thread. This simulates what you might get if you were to read from an active data source.

```
def load_data(q):
    i = 0
    while True:
        q.put(i)
        sleep(random())
        i += 1

>>> from threading import Thread
>>> load_thread = Thread(target=load_data, args=(input_q,))
>>> load_thread.start()

>>> result_q.qsize()
4
>>> result_q.qsize()
9
```

We consume data from the result_q and print results to the screen.

```
>>> while True:
... item = result_q.get()
... print(item)
2
4
6
8
10
12
...
```

3.13.2 Limitations

• This doesn't do any sort of auto-batching of computations, so ideally you batch your data to take significantly longer than 1ms to run.

• This isn't a proper streaming system. There is no support outside of what you see here. In particular there are no policies for dropping data, joining over time windows, etc..

3.13.3 Extensions

We can extend this small example to more complex systems that have buffers, split queues, merge queues, etc. all by manipulating normal Python Queues.

Here are a couple of useful function to multiplex and merge queues:

```
from queue import Queue
from threading import Thread
def multiplex(n, q, **kwargs):
    """ Convert one queue into several equivalent Queues
    >>> q1, q2, q3 = multiplex(3, in_q)
   out_queues = [Queue(**kwargs) for i in range(n)]
    def f():
        while True:
            x = q.get()
            for out_q in out_queues:
               out_q.put(x)
   t = Thread(target=f)
   t.daemon = True
   t.start()
    return out_queues
def push(in_q, out_q):
    while True:
        x = in_q.get()
        out_q.put(x)
def merge(*in_qs, **kwargs):
    """ Merge multiple queues together
    >>> out_q = merge(q1, q2, q3)
   out_q = Queue(**kwargs)
   threads = [Thread(target=push, args=(q, out_q)) for q in in_qs]
    for t in threads:
        t.daemon = True
        t.start()
    return out_q
```

With useful functions like these we can build out more sophisticated data processing pipelines that split off and join back together. By creating queues with maxsize= we can control buffering and apply back pressure.

3.14 API

Executor

Executor([address, start, loop, timeout,])	Drive computations on a distributed cluster
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Executor.cancel(futures)	Cancel running futures
Executor.compute(args[, sync])	Compute dask collections on cluster
Executor.gather(futures[, errors, maxsize])	Gather futures from distributed memory
Executor.get(dsk, keys[, restrictions,])	Compute dask graph
Executor.has_what([workers])	Which keys are held by which workers
Executor.map(func, *iterables, **kwargs)	Map a function on a sequence of arguments
Executor.ncores([workers])	The number of threads/cores available on each worker node
Executor.persist(collections)	Persist dask collections on cluster
Executor.rebalance([futures, workers])	Rebalance data within network
<pre>Executor.replicate(futures[, n, workers,])</pre>	Set replication of futures within network
Executor.restart()	Restart the distributed network
<pre>Executor.run(function, *args, **kwargs)</pre>	Run a function on all workers outside of task scheduling system
<pre>Executor.scatter(data[, workers, broadcast,])</pre>	Scatter data into distributed memory
Executor.shutdown([timeout])	Send shutdown signal and wait until scheduler terminates
Executor.submit(func, *args, **kwargs)	Submit a function application to the scheduler
Executor.upload_file(filename)	Upload local package to workers
Executor.who_has([futures])	The workers storing each future's data

Future

Future(key, executor)	A remotely running computation
Future.cancel()	Returns True if the future has been cancelled
Future.cancelled()	Returns True if the future has been cancelled
Future.done()	Is the computation complete?
Future.exception()	Return the exception of a failed task
Future.result()	Wait until computation completes.
Future.traceback()	Return the traceback of a failed task

Other

as_completed(fs)	Return futures in the order in which they complete
distributed.diagnostics.progress(*futures,)	Track progress of futures
wait(fs[, timeout, return_when])	Wait until all futures are complete

3.14.1 Executor

 $\begin{array}{ll} \textbf{class} \ \texttt{distributed.executor.Executor} \ (address=None, & \textit{start=True}, & loop=None, & \textit{timeout=3}, \\ & \textit{set_as_default=False}) \end{array}$

Drive computations on a distributed cluster

The Executor connects users to a distributed compute cluster. It provides an asynchronous user interface around functions and futures. This class resembles executors in concurrent.futures but also allows Future objects within submit/map calls.

Parameters address: string, tuple, or "Scheduler"

This can be the address of a Scheduler server, either as a string '127.0.0.1:8787' or tuple ('127.0.0.1', 8787) or it can be a local Scheduler object.

See also:

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distributed.scheduler.Scheduler Internal scheduler

Examples

Provide cluster's head node address on initialization:

```
>>> executor = Executor('127.0.0.1:8787')
```

Use submit method to send individual computations to the cluster

```
>>> a = executor.submit(add, 1, 2)
>>> b = executor.submit(add, 10, 20)
```

Continue using submit or map on results to build up larger computations

```
>>> c = executor.submit(add, a, b)
```

Gather results with the gather method.

```
>>> executor.gather([c])
33
```

cancel (futures)

Cancel running futures

This stops future tasks from being scheduled if they have not yet run and deletes them if they have already run. After calling, this result and all dependent results will no longer be accessible

Parameters futures: list of Futures

```
compute (args, sync=False)
```

Compute dask collections on cluster

Parameters args: iterable of dask objects or single dask object

Collections like dask.array or dataframe or dask.value objects

```
sync: bool (optional)
```

Returns Futures if False (default) or concrete values if True

Returns List of Futures if input is a sequence, or a single future otherwise

See also:

Executor. **get** Normal synchronous dask.get function

Examples

```
>>> from dask import do, value
>>> from operator import add
>>> x = dask.do(add)(1, 2)
>>> y = dask.do(add)(x, x)
>>> xx, yy = executor.compute([x, y])
>>> xx
<fruture: status: finished, key: add-8f6e709446674bad78ea8aeecfee188e>
>>> xx.result()
3
>>> yy.result()
6
```

Also support single arguments

```
>>> xx = executor.compute(x)
```

```
gather (futures, errors='raise', maxsize=0)
```

Gather futures from distributed memory

Accepts a future, nested container of futures, iterator, or queue. The return type will match the input type.

Returns Future results

See also:

Executor.scatter Send data out to cluster

Examples

```
>>> from operator import add
>>> e = Executor('127.0.0.1:8787')
>>> x = e.submit(add, 1, 2)
>>> e.gather(x)
3
>>> e.gather([x, [x], x]) # support lists and dicts
[3, [3], 3]
```

```
>>> seq = e.gather(iter([x, x])) # support iterators
>>> next(seq)
3
```

get (dsk, keys, restrictions=None, loose_restrictions=None)

Compute dask graph

Parameters dsk: dict

keys: object, or nested lists of objects

restrictions: dict (optional)

A mapping of {key: {set of worker hostnames}} that restricts where jobs can take place

See also:

Executor. compute Compute asynchronous collections

Examples

```
>>> from operator import add
>>> e = Executor('127.0.0.1:8787')
>>> e.get({'x': (add, 1, 2)}, 'x')
3
```

has_what (workers=None)

Which keys are held by which workers

Parameters workers: list (optional)

A list of worker addresses, defaults to all

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See also:

```
Executor.who_has, Executor.ncores
```

Examples

```
map (func, *iterables, **kwargs)
```

Map a function on a sequence of arguments

Arguments can be normal objects or Futures

Parameters func: callable

iterables: Iterables, Iterators, or Queues

pure: bool (defaults to True)

Whether or not the function is pure. Set pure=False for impure functions like np.random.random.

workers: set, iterable of sets

A set of worker hostnames on which computations may be performed. Leave empty to default to all workers (common case)

Returns List, iterator, or Queue of futures, depending on the type of the inputs.

See also:

Executor. submit Submit a single function

Examples

```
>>> L = executor.map(func, sequence)
```

```
nbytes (keys=None, summary=True)
```

The bytes taken up by each key on the cluster

This is as measured by sys.getsizeof which may not accurately reflect the true cost.

Parameters keys: list (optional)

A list of keys, defaults to all keys

summary: boolean, (optional)

Summarize keys into key types

See also:

Executor.who_has

Examples

```
>>> x, y, z = e.map(inc, [1, 2, 3])
>>> e.nbytes(summary=False)
{'inc-1c8dd6be1c21646c71f76c16d09304ea': 28,
    'inc-1e297fc27658d7b67b3a758f16bcf47a': 28,
    'inc-fd65c238a7ea60f6a01bf4c8a5fcf44b': 28}
```

```
>>> e.nbytes(summary=True)
{'inc': 84}
```

ncores (workers=None)

The number of threads/cores available on each worker node

Parameters workers: list (optional)

A list of workers that we care about specifically. Leave empty to receive information about all workers.

See also:

```
Executor.who_has, Executor.has_what
```

Examples

```
>>> e.ncores()
{'192.168.1.141:46784': 8,
    '192.167.1.142:47548': 8,
    '192.167.1.143:47329': 8,
    '192.167.1.144:37297': 8}
```

persist (collections)

Persist dask collections on cluster

Starts computation of the collection on the cluster in the background. Provides a new dask collection that is semantically identical to the previous one, but now based off of futures currently in execution.

Parameters collections: sequence or single dask object

Collections like dask.array or dataframe or dask.value objects

Returns List of collections, or single collection, depending on type of input.

See also:

```
Executor.compute
```

Examples

```
>>> xx = executor.persist(x)
>>> xx, yy = executor.persist([x, y])
```

processing(workers=None)

The tasks currently running on each worker

Parameters workers: list (optional)

A list of worker addresses, defaults to all

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See also:

```
Executor.stacks, Executor.who_has, Executor.has_what, Executor.ncores
```

Examples

rebalance (futures=None, workers=None)

Rebalance data within network

Move data between workers to roughly balance memory burden. This either affects a subset of the keys/workers or the entire network, depending on keyword arguments.

This operation is generally not well tested against normal operation of the scheduler. It it not recommended to use it while waiting on computations.

Parameters futures: list, optional

A list of futures to balance, defaults all data

workers: list, optional

A list of workers on which to balance, defaults to all workers

```
replicate (futures, n=None, workers=None, branching_factor=2)
```

Set replication of futures within network

This performs a tree copy of the data throughout the network individually on each piece of data.

This operation blocks until complete. It does not guarantee replication of data to future workers.

Parameters futures: list of futures

Futures we wish to replicate

n: int, optional

Number of processes on the cluster on which to replicate the data. Defaults to all.

workers: list of worker addresses

Workers on which we want to restrict the replication. Defaults to all.

branching_factor: int, optional

The number of workers that can copy data in each generation

See also:

```
Executor.rebalance
```

Examples

```
>>> x = e.submit(func, *args)
>>> e.replicate([x]) # send to all workers
>>> e.replicate([x], n=3) # send to three workers
>>> e.replicate([x], workers=['alice', 'bob']) # send to specific
```

```
>>> e.replicate([x], n=1, workers=['alice', 'bob']) # send to one of specific
>>> e.replicate([x], n=1) # reduce replications
```

restart()

Restart the distributed network

This kills all active work, deletes all data on the network, and restarts the worker processes.

```
run (function, *args, **kwargs)
```

Run a function on all workers outside of task scheduling system

This calls a function on all currently known workers immediately, blocks until those results come back, and returns the results asynchronously as a dictionary keyed by worker address. This method if generally used for side effects, such and collecting diagnostic information or installing libraries.

Parameters function: callable

*args: arguments for remote function

**kwargs: keyword arguments for remote function

workers: list

Workers on which to run the function. Defaults to all known workers.

Examples

```
>>> e.run(os.getpid)
{'192.168.0.100:9000': 1234,
    '192.168.0.101:9000': 4321,
    '192.168.0.102:9000': 5555}
```

Restrict computation to particular workers with the workers = keyword argument.

```
>>> e.run(os.getpid, workers=['192.168.0.100:9000',
...
'192.168.0.101:9000'])
{'192.168.0.101:9000': 1234,
'192.168.0.101:9000': 4321}
```

scatter (data, workers=None, broadcast=False, maxsize=0)

Scatter data into distributed memory

Parameters data: list, iterator, dict, or Queue

Data to scatter out to workers. Output type matches input type.

workers: list of tuples (optional)

Optionally constrain locations of data. Specify workers as hostname/port pairs, e.g. ('127.0.0.1', 8787).

broadcast: bool (defaults to False)

Whether to send each data element to all workers. By default we round-robin based on number of cores.

maxsize: int (optional)

Maximum size of queue if using queues, 0 implies infinite

Returns List, dict, iterator, or queue of futures matching the type of input.

See also:

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Executor. gather Gather data back to local process

Examples

```
>>> e.scatter({'x': 1, 'y': 2, 'z': 3})
{'x': <Future: status: finished, key: x>,
   'y': <Future: status: finished, key: y>,
   'z': <Future: status: finished, key: z>}
```

Constrain location of data to subset of workers >>> e.scatter([1, 2, 3], workers=[('hostname', 8788)]) # doctest: +SKIP

Handle streaming sequences of data with iterators or queues >>> seq = e.scatter(iter([1, 2, 3])) # doctest: +SKIP >>> next(seq) # doctest: +SKIP <Future: status: finished, key: c0a8a20f903a4915b94db8de3ea63195>,

Broadcast data to all workers >>> [future] = e.scatter([element], broadcast=True) # doctest: +SKIP

```
shutdown (timeout=10)
```

Send shutdown signal and wait until scheduler terminates

```
stacks (workers=None)
```

The task queues on each worker

Parameters workers: list (optional)

A list of worker addresses, defaults to all

See also:

Executor.processing, Executor.who_has, Executor.has_what, Executor.ncores

Examples

```
start (**kwargs)
```

Start scheduler running in separate thread

```
submit (func, *args, **kwargs)
```

Submit a function application to the scheduler

Parameters func: callable

*args:

**kwargs:

pure: bool (defaults to True)

Whether or not the function is pure. Set pure=False for impure functions like np.random.random.

workers: set, iterable of sets

A set of worker hostnames on which computations may be performed. Leave empty to default to all workers (common case)

allow_other_workers: bool (defaults to False)

Used with *workers*. Inidicates whether or not the computations may be performed on workers that are not in the *workers* set(s).

Returns Future

See also:

Executor.map Submit on many arguments at once

Examples

```
>>> c = executor.submit(add, a, b)
```

upload_file (filename)

Upload local package to workers

This sends a local file up to all worker nodes. This file is placed into a temporary directory on Python's system path so any .py or .egg files will be importable.

Parameters filename: string

Filename of .py or .egg file to send to workers

Examples

```
>>> executor.upload_file('mylibrary.egg')
>>> from mylibrary import myfunc
>>> L = e.map(myfunc, seq)
```

who_has (futures=None)

The workers storing each future's data

Parameters futures: list (optional)

A list of futures, defaults to all data

See also:

```
Executor.has_what, Executor.ncores
```

Examples

```
>>> x, y, z = e.map(inc, [1, 2, 3])
>>> wait([x, y, z])
>>> e.who_has()
{'inc-1c8dd6be1c21646c71f76c16d09304ea': ['192.168.1.141:46784'],
    'inc-1e297fc27658d7b67b3a758f16bcf47a': ['192.168.1.141:46784'],
    'inc-fd65c238a7ea60f6a01bf4c8a5fcf44b': ['192.168.1.141:46784']}
```

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3.14.2 CompatibleExecutor

class distributed.executor.CompatibleExecutor (address=None, start=True, loop=None, time-out=3, $set_as_default=False$)

A concurrent.futures-compatible Executor

A subclass of Executor that conforms to concurrent futures API, allowing swapping in for other Executors.

```
map (func, *iterables, **kwargs)
```

Map a function on a sequence of arguments

Returns iter results: iterable

Iterable yielding results of the map.

See also:

Executor.map for more info

3.14.3 Future

class distributed.executor.Future (key, executor)

A remotely running computation

A Future is a local proxy to a result running on a remote worker. A user manages future objects in the local Python process to determine what happens in the larger cluster.

See also:

Executor Creates futures

Examples

Futures typically emerge from Executor computations

```
>>> my_future = executor.submit(add, 1, 2)
```

We can track the progress and results of a future

```
>>> my_future <Future: status: finished, key: add-8f6e709446674bad78ea8aeecfee188e>
```

We can get the result or the exception and traceback from the future

```
>>> my_future.result()
```

cancel()

Returns True if the future has been cancelled

cancelled()

Returns True if the future has been cancelled

```
done()
    Is the computation complete?

exception()
    Return the exception of a failed task
    See also:
    Future.traceback

result()
    Wait until computation completes. Gather result to local process
```

wait until computation completes.

traceback()

Return the traceback of a failed task

This returns a traceback object. You can inspect this object using the traceback module. Alternatively if you call future.result() this traceback will accompany the raised exception.

See also:

Future.exception

Examples

```
>>> import traceback
>>> tb = future.traceback()
>>> traceback.export_tb(tb)
[...]
```

3.14.4 Other

```
distributed.executor.as\_completed(fs)
```

Return futures in the order in which they complete

This returns an iterator that yields the input future objects in the order in which they complete. Calling next on the iterator will block until the next future completes, irrespective of order.

This function does not return futures in the order in which they are input.

```
distributed.diagnostics.progress(*futures, **kwargs)
    Track progress of futures
```

This operates differently in the notebook and the console

- •Notebook: This returns immediately, leaving an IPython widget on screen
- •Console: This blocks until the computation completes

Parameters futures: Futures

A list of futures or keys to track

notebook: bool (optional)

Running in the notebook or not (defaults to guess)

multi: bool (optional)

Track different functions independently (defaults to True)

complete: bool (optional)

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Track all keys (True) or only keys that have not yet run (False) (defaults to True)

Examples

```
>>> progress(futures)
[###########################| | 100% Completed | 1.7s
```

distributed.executor.wait (fs, timeout=None, return_when='ALL_COMPLETED')

Wait until all futures are complete

Parameters fs: list of futures

Returns Named tuple of completed, not completed

3.15 Foundations

You should read through the quickstart before reading this document.

Distributed computing is hard for two reasons:

- 1. Consistent coordination of distributed systems requires sophistication
- 2. Concurrent network programming is tricky and error prone

The foundations of distributed provide abstractions to hide some complexity of concurrent network programming (#2). These abstractions ease the construction of sophisticated parallel systems (#1) in a safer environment. However, as with all layered abstractions, ours has flaws. Critical feedback is welcome.

3.15.1 Concurrency with Tornado Coroutines

Worker and Scheduler nodes operate concurrently. They serve several overlapping requests and perform several overlapping computations at the same time without blocking. There are several approaches for concurrent programming, we've chosen to use Tornado for the following reasons:

- 1. Developing and debugging is more comfortable without threads
- 2. Tornado's documentation is excellent
- 3. Stackoverflow coverage is excellent
- 4. Performance is satisfactory

3.15.2 Communication with Tornado Streams (raw sockets)

Workers, the Scheduler, and clients communicate with each other over the network. They use *raw sockets* as mediated by tornado streams. We separate messages by a sentinel value.

```
distributed.core.read(stream)
Read a message from a stream
distributed.core.write(stream, msg)
Write a message to a stream
```

3.15.3 Servers

Worker and Scheduler nodes serve requests over TCP. Both Worker and Scheduler objects inherit from a Server class. This Server class thinly wraps tornado.tcpserver.TCPServer. These servers expect requests of a particular form.

Superclass for both Worker and Center objects. Inherits from tornado.tcpserver.TCPServer, adding a protocol for RPC.

Handlers

Servers define operations with a handlers dict mapping operation names to functions. The first argument of a handler function must be a stream for the connection to the client. Other arguments will receive inputs from the keys of the incoming message which will always be a dictionary.

```
>>> def pingpong(stream):
... return b'pong'
```

```
>>> def add(stream, x, y):
... return x + y
```

```
>>> handlers = {'ping': pingpong, 'add': add}
>>> server = Server(handlers)
>>> server.listen(8000)
```

Message Format

The server expects messages to be dictionaries with a special key, 'op' that corresponds to the name of the operation, and other key-value pairs as required by the function.

So in the example above the following would be good messages.

```
•{'op': 'ping'}
•{'op': 'add': 'x': 10, 'y': 20}
```

3.15.4 RPC

To interact with remote servers we typically use rpc objects.

Normally we construct messages as dictionaries and send them with read/write

```
>>> stream = yield connect(ip, port)
>>> msg = {'op': 'add', 'x': 10, 'y': 20}
>>> yield write(stream, msg)
>>> response = yield read(stream)
```

To reduce verbosity we use an rpc object.

```
>>> remote = rpc(ip=ip, port=port)
>>> response = yield remote.add(x=10, y=20)
```

One rpc object can be reused for several interactions. Additionally, this object creates and destroys many streams as necessary and so is safe to use in multiple overlapping communications.

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When done, close streams explicitly.

```
>>> remote.close_streams()
```

3.15.5 Example

Here is a small example using distributed.core to create and interact with a custom server.

Server Side

```
from tornado import gen
from tornado.ioloop import IOLoop
from distributed.core import write, Server

def add(stream, x=None, y=None): # simple handler, just a function
    return x + y

@gen.coroutine
def stream_data(stream, interval=1): # complex handler, multiple responses
    data = 0
    while True:
        yield gen.sleep(interval)
        data += 1
        yield write(stream, data)

s = Server({'add': add, 'stream': stream_data})
s.listen(8888)

IOLoop.current().start()
```

Client Side

```
from tornado import gen
from tornado.ioloop import IOLoop
from distributed.core import connect, read, write
@gen.coroutine
def f():
   stream = yield connect('127.0.0.1', 8888)
   yield write(stream, {'op': 'add', 'x': 1, 'y': 2})
   result = yield read(stream)
   print (result)
>>> IOLoop().run_sync(f)
@gen.coroutine
def g():
    stream = yield connect('127.0.0.1', 8888)
   yield write(stream, {'op': 'stream', 'interval': 1})
    while True:
       result = yield read(stream)
        print (result)
```

```
>>> IOLoop().run_sync(g)
1
2
3
...
```

Client Side with rpc

RPC provides a more pythonic interface. It also provides other benefits, such as using multiple streams in concurrent cases. Most distributed code uses rpc. The exception is when we need to perform multiple reads or writes, as with the stream data case above.

```
from tornado import gen
from tornado.ioloop import IOLoop
from distributed.core import rpc

@gen.coroutine
def f():
    # stream = yield connect('127.0.0.1', 8888)
    # yield write(stream, {'op': 'add', 'x': 1, 'y': 2})
    # result = yield read(stream)
    r = rpc(ip='127.0.0.1', 8888)
    result = yield r.add(x=1, y=2)

    print(result)

>>> IOLoop().run_sync(f)
3
```

3.15.6 Everything is a Server

Workers, Scheduler, and Nanny objects all inherit from Server. Each maintains separate state and serves separate functions but all communicate in the way shown above. They talk to each other by opening connections, writing messages that trigger remote functions, and then collect the results with read.

3.16 Client Interaction

As discussed in the quickstart users can interact with the worker-center network with the Executor abstraction.

This is built with lower level functions described below.

3.16.1 Scatter/Gather

Users rarely create RemoteData objects by hand. They are created by other client libraries or functions like gather and scatter.

```
distributed.client.scatter(center, data, serialize=True)
Scatter data to workers
```

Parameters center:

(ip, port) tuple or Stream, or rpc object designating the Center

data: dict or iterable

either a dictionary of key: value pairs or an iterable of values

key:

if data is an iterable of values then we use the key to generate keys as key-0, key-1, key-2, ...

See also:

```
distributed.client.gather,
distributed.client.scatter to workers
```

Examples

```
>>> remote_data = scatter('127.0.0.1:8787', [1, 2, 3])
>>> local_data = gather('127.0.0.1:8787', remote_data)
```

```
distributed.client.gather(center, needed)
```

Gather data from distributed workers

This operates by first asking the center who has all of the state keys and then trying those workers directly.

Keys not found on the network will not appear in the output. You should check the length of the output against the input if concerned about missing data.

Parameters center:

(ip, port) tuple or Stream, or rpc object designating the Center

needed: iterable

A list of required keys

Returns result: dict

A mapping of the given keys to data values

See also:

```
distributed.client.scatter,
distributed.client.gather_from_workers
```

Examples

```
>>> remote_data = scatter('127.0.0.1:8787', [1, 2, 3])
>>> local_data = gather('127.0.0.1:8787', remote_data)
```

```
distributed.client.delete(center, keys)
```

Delete keys from all workers

```
distributed.client.clear(center)
```

Clear all data from all workers' memory

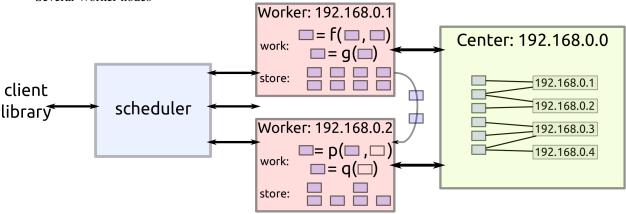
See also:

```
distributed.client.delete
```

3.17 Worker

We build a distributed network from two kinds of nodes.

- A single scheduler node
- · Several Worker nodes



This page describes the worker nodes.

3.17.1 Serve Data

Workers serve data from a local dictionary of data:

```
{'x': np.array(...),
    'y': pd.DataFrame(...)}
```

Operations include normal dictionary operations, like get, set, and delete key-value pairs. In the following example we connect to two workers, collect data from one worker and send it to another.

```
alice = rpc(ip='192.168.0.101', port=8788)
d = yield alice.get_data(keys=['x', 'y'])
bob = rpc(ip='192.168.0.102', port=8788)
yield bob.update_data(data=d)
```

However, this is only an example, typically one does not manually manage data transfer between workers. They handle that as necessary on their own.

3.17.2 Compute

Workers evaluate functions provided by the user on their data. They evaluate functions either on their data or can automatically collect data from peers (as shown above) if they don't have the necessary data but their peers do:

```
z \leftarrow add(x, y) # can be done with only local data z \leftarrow add(x, a) # need to find out where we can get 'a'
```

The result of such a computation on our end is just a response b'OK'. The actual result stays on the remote worker.

```
>>> response, metadata = yield alice.compute(function=add, keys=['x', 'a'])
>>> response
b'OK'
```

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```
>>> metadata {'nbytes': 1024}
```

The worker also reports back to the center/scheduler whenever it completes a computation. Metadata storage is centralized but all data transfer is peer-to-peer. Here is a quick example of what happens during a call to compute:

```
client: Hey Alice! Compute ``z <- add(x, a)``
Alice: Hey Center! Who has a?
Center: Hey Alice! Bob has a.
Alice: Hey Bob! Send me a!
Bob: Hey Alice! Here's a!

Alice: Hey Client! I've computed z and am holding on to it!
Alice: Hey Center! I have z!</pre>
```

Worker Node

Workers perform two functions:

- 1.Serve data from a local dictionary
- 2.**Perform computation** on that data and on data from peers

Additionally workers keep a scheduler informed of their data and use that scheduler to gather data from other workers when necessary to perform a computation.

You can start a worker with the dworker command line application:

```
$ dworker scheduler-ip:port
```

State

```
•data: {key: object}: Dictionary mapping keys to actual values
•active: {key}: Set of keys currently under computation
•ncores: int: Number of cores used by this worker process
•executor: concurrent.futures.ThreadPoolExecutor: Executor used to perform computation
•local_dir: path: Path on local machine to store temporary files
•scheduler: rpc: Location of scheduler. See .ip/.port attributes.
•name: string: Alias
•services: {str: Server}: Auxiliary web servers running on this worker
•service_ports: {str: port}:

See also:
```

Examples

Create schedulers and workers in Python:

distributed.scheduler.Scheduler

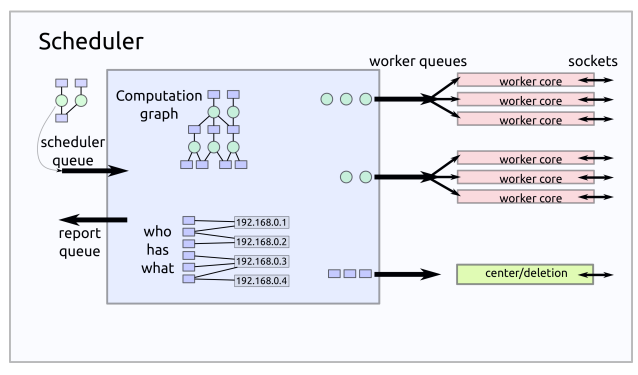
```
>>> from distributed import Scheduler, Worker
>>> c = Scheduler('192.168.0.100', 8787)
>>> w = Worker(c.ip, c.port)
>>> yield w._start(port=8788)
```

Or use the command line:

```
$ dask-scheduler
Start scheduler at 127.0.0.1:8787

$ dask-worker 127.0.0.1:8787
Start worker at: 127.0.0.1:8788
Registered with scheduler at: 127.0.0.1:8787
```

3.18 Scheduler



The scheduler orchestrates which workers work on which tasks in what order. It tracks the current state of the entire cluster. It consists of several coroutines rounning in a single event loop.

Dynamic distributed task scheduler

The scheduler tracks the current state of workers, data, and computations. The scheduler listens for events and responds by controlling workers appropriately. It continuously tries to use the workers to execute an ever growing dask graph.

All events are handled quickly, in linear time with respect to their input (which is often of constant size) and generally within a millisecond. To accomplish this the scheduler tracks a lot of state. Every operation maintains the consistency of this state.

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The scheduler communicates with the outside world either by adding pairs of in/out queues or by responding to a new IOStream (the Scheduler can operate as a typical distributed Server). It maintains a consistent and valid view of the world even when listening to several clients at once.

A Scheduler is typically started either with the dask-scheduler executable:

```
$ dask-scheduler
```

Or as part of when an Executor starts up and connects to a Center:

```
>>> e = Executor('127.0.0.1:8787')
>>> e.scheduler
Scheduler(...)
```

Users typically do not interact with the scheduler except through Plugins. See http://distributed.readthedocs.io/en/latest/plugins.html

State

```
•tasks: {key: task}: Dictionary mapping key to task, either dask task, or serialized dict like:
     {'function': b'xxx', 'args': b'xxx'} or {'task': b'xxx'}
•dependencies: {key: {key}}: Dictionary showing which keys depend on which others
•dependents: {key: {key}}: Dictionary showing which keys are dependent on which others
•waiting: {key}}: Dictionary like dependencies but excludes keys already computed
•waiting_data: {key: {key}}: Dictionary like dependents but excludes keys already computed
•ready: deque (key) Keys that are ready to run, but not yet assigned to a worker
•ncores: {worker: int}: Number of cores owned by each worker
•idle: {worker}: Set of workers that are not fully utilized
•services: {str: port}: Other services running on this scheduler, like HTTP
•worker_info: {worker: {str: data}}: Information about each worker
•host_info: {hostname: dict}: Information about each worker host
•who_has: {key: {worker}}: Where each key lives. The current state of distributed memory.
•has what: {worker:
                       {key}}: What worker has what keys. The transpose of who_has.
•who_wants: {key:
                     {client}}: Which clients want each key. The active targets of computation.
•wants_what: {client: {key}}: What keys are wanted by each client.. The transpose of
    who wants.
•nbytes: {key: int}: Number of bytes for a key as reported by workers holding that key.
•processing: {worker: {key: cost}}: Set of keys currently in execution on each worker and
    their expected duration
•rprocessing: {key: {worker}}: Set of workers currently executing a particular task
•task_duration: {key-prefix: time} Time we expect certain functions to take, e.g. {'sum':
     0.25}
•occupancy: {worker: time} Expected runtime for all tasks currently processing on a worker
•stacks: {worker: [keys]}: List of keys waiting to be sent to each worker
•released: {keys} Set of keys that are known, but released from memory
•unrunnable: {key} Keys that we are unable to run
```

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```
•retrictions: {key:
                          {hostnames}}: A set of hostnames per key of where that key can be run. Usu-
         ally this is empty unless a key has been specifically restricted to only run on certain hosts. These
         restrictions don't include a worker port. Any worker on that hostname is deemed valid.
    •loose_retrictions: {key}: Set of keys for which we are allow to violate restrictions (see above) if not
         valid workers are present.
    •suspicious tasks: {key: int} Number of times a task has been involved in a worker failure
    •keyorder: {key: tuple}: A score per key that determines its priority
    •scheduler_queues: [Queues]: A list of Tornado Queues from which we accept stimuli
    •report_queues: [Queues]: A list of Tornado Queues on which we report results
    •streams: [IOStreams]: A list of Tornado IOStreams from which we both accept stimuli and report
         results
    •coroutines: [Futures]: A list of active futures that control operation
    •exceptions: {key: Exception}: A dict mapping keys to remote exceptions
    •tracebacks: {key: list}: A dict mapping keys to remote tracebacks stored as a list of strings
    •exceptions blame: {key: key}: A dict mapping a key to another key on which it depends that has
         failed
    •deleted_keys: {key: {workers}} Locations of workers that have keys that should be deleted
    •loop: IOLoop: The running Torando IOLoop
add client(stream, client=None)
     Listen to messages from an IOStream
add_plugin (plugin)
     Add external plugin to scheduler
     See http://http://distributed.readthedocs.io/en/latest/plugins.html
broadcast (stream=None, msg=None, workers=None)
     Broadcast message to workers, return all results
cancel (stream, keys=None, client=None)
     Stop execution on a list of keys
cleanup()
     Clean up queues and coroutines, prepare to stop
clear_data_from_workers()
     This is intended to be run periodically,
     The self._delete_periodic_callback attribute holds a PeriodicCallback that runs this every
     self.delete interval milliseconds".
close (stream=None, fast=False)
     Send cleanup signal to all coroutines then wait until finished
     See also:
     Scheduler.cleanup
coerce_address(addr)
```

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Coerce possible input addresses to canonical form

Handles lists, strings, bytes, tuples, or aliases

correct time delay (worker, msg)

Apply offset time delay in message times

Operates in place

ensure_idle_ready()

Run ready tasks on idle workers

Work stealing policy

If some workers are idle but not others, if there are no globally ready tasks, and if there are tasks in worker stacks, then we start to pull preferred tasks from overburdened workers and deploy them back into the global pool in the following manner.

We determine the number of tasks to reclaim as the number of all tasks in all stacks times the fraction of idle workers to all workers. We sort the stacks by size and walk through them, reclaiming half of each stack until we have enough task to fill the global pool. We are careful not to reclaim tasks that are restricted to run on certain workers.

ensure_in_play(key)

Ensure that a key is on track to enter memory in the future

This will only act on keys currently in self.released.

ensure_occupied_stacks(worker)

Send tasks to worker while it has tasks and free cores

These tasks may come from the worker's own stacks or from the global ready deque.

We update the idle workers set appropriately.

finished()

Wait until all coroutines have ceased

forget (key)

Forget a key if no one cares about it

This removes all knowledge of how to produce a key from the scheduler. This is almost exclusively called by release_held_data

gather (stream=None, keys=None)

Collect data in from workers

handle_messages (in_queue, report, client=None)

Master coroutine. Handles inbound messages.

This runs once per Queue or Stream.

handle queues (scheduler queue, report queue)

Register new control and report queues to the Scheduler

identity(stream)

Basic information about ourselves and our cluster

issaturated(worker, latency=0.005)

A worker is saturated if it has enough work to avoid being idle

A worker is saturated if the following criteria are met

- 1.It is working on at least as many tasks as it has cores
- 2. The expected time it will take to complete all of its currently assigned tasks is at least a full round-trip time. This is relevant when it has many small tasks

```
log state(msg='')
    Log current full state of the scheduler
mark_failed (key, failing_key=None)
     When a task fails mark it and all dependent task as failed
mark_key_in_memory (key, workers=None, type=None)
     Mark that a key now lives in distributed memory
mark_missing_data(keys=None, key=None, worker=None, **kwargs)
     Mark that certain keys have gone missing. Recover.
     See also:
     recover missing
mark_not_processing(key, worker)
     Mark that a key is done running on a worker
mark_processing (key, worker, latency=5e-05)
     Mark that a key is running on a worker
mark_ready_to_run(key)
     Mark a task as ready to run.
     If the task should be assigned to a worker then make that determination and assign appropriately. Otherwise
     place task in the ready queue.
     Trigger appropriate workers if idle.
     See also:
     decide_worker, Scheduler.ensure_occupied
mark_task_erred (key=None, worker=None, exception=None, traceback=None, **kwargs)
     Mark that a task has erred on a particular worker
     See also:
     Scheduler.mark_failed
mark_task_finished (key=None, worker=None, nbytes=None, type=None, compute_start=None,
                         compute stop=None, transfer start=None, transfer stop=None, **kwargs)
     Mark that a task has finished execution on a particular worker
mark_task_killed_worker(key=None, worker=None)
     Mark that is likely killing workers
     Place a message into the scheduler's queue
recover_missing(key)
     Recover a recently lost piece of data
     This assumes that we've already removed this key from who_has/has_what.
release_held_data(keys=None)
```

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Mark that a key is no longer externally required to be in memory

We no longer need to keep data in memory to compute this This occurs after we've computed it or after we've forgotten it

release_live_dependencies (key)

```
remove worker (stream=None, address=None)
     Mark that a worker no longer seems responsive
     See also:
     Scheduler.recover_missing
replicate (stream=None, keys=None, n=None, workers=None, branching factor=2)
     Replicate data throughout cluster
     This performs a tree copy of the data throughout the network individually on each piece of data.
         Parameters keys: Iterable
               list of keys to replicate
             n: int
               Number of replications we expect to see within the cluster
             branching_factor: int, optional
               The number of workers that can copy data in each generation
     See also:
     Scheduler.rebalance
report (msg)
     Publish updates to all listening Queues and Streams
restart()
     Restart all workers. Reset local state
scatter(stream=None, data=None, workers=None, client=None, broadcast=False)
     Send data out to workers
should_steal (key, bandwidth=None)
     Is a key good for stealing from its chosen worker?
     It must have the following attributes
       1.Not have too many dependencies
     2. Not be restricted to run on that worker 3 Take less time to transfer than to compute
start (port=8786, start queues=True)
     Clear out old state and restart all running coroutines
update_data(who_has=None, nbytes=None, client=None)
     Learn that new data has entered the network from an external source
     See also:
     Scheduler.mark_key_in_memory
update_graph (client=None, tasks=None, keys=None, dependencies=None, restrictions=None,
                  loose_restrictions=None)
     Add new computations to the internal dask graph
     This happens whenever the Executor calls submit, map, get, or compute.
workers_list (workers)
     List of qualifying workers
```

Takes a list of worker addresses or hostnames. Returns a list of all worker addresses that match

distributed.scheduler.decide_worker(dependencies, stacks, processing, who_has, has_what, restrictions, loose restrictions, nbytes, key)

Decide which worker should take task

```
>>> dependencies = {'c': {'b'}, 'b': {'a'}}
>>> stacks = {'alice:8000': ['z'], 'bob:8000': []}
>>> processing = {'alice:8000': set(), 'bob:8000': set()}
>>> who_has = {'a': {'alice:8000'}}
>>> has_what = {'alice:8000': {'a'}}
>>> nbytes = {'a': 100}
>>> restrictions = {}
>>> loose_restrictions = set()
```

We choose the worker that has the data on which 'b' depends (alice has 'a')

```
>>> decide_worker(dependencies, stacks, processing, who_has, has_what,
... restrictions, loose_restrictions, nbytes, 'b')
'alice:8000'
```

If both Alice and Bob have dependencies then we choose the less-busy worker

```
>>> who_has = {'a': {'alice:8000', 'bob:8000'}}
>>> has_what = {'alice:8000': {'a'}, 'bob:8000': {'a'}}
>>> decide_worker(dependencies, stacks, processing, who_has, has_what,
... restrictions, loose_restrictions, nbytes, 'b')
'bob:8000'
```

Optionally provide restrictions of where jobs are allowed to occur

```
>>> restrictions = {'b': {'alice', 'charlie'}}
>>> decide_worker(dependencies, stacks, processing, who_has, has_what,
... restrictions, loose_restrictions, nbytes, 'b')
'alice:8000'
```

If the task requires data communication, then we choose to minimize the number of bytes sent between workers. This takes precedence over worker occupancy.

```
>>> dependencies = {'c': {'a', 'b'}}
>>> who_has = {'a': {'alice:8000'}, 'b': {'bob:8000'}}
>>> has_what = {'alice:8000': {'a'}, 'bob:8000': {'b'}}
>>> nbytes = {'a': 1, 'b': 1000}
>>> stacks = {'alice:8000': [], 'bob:8000': []}
```

```
>>> decide_worker(dependencies, stacks, processing, who_has, has_what,
... {}, set(), nbytes, 'c')
'bob:8000'
```

3.19 Resilience

Software fails, Hardware fails, network connections fail, user code fails. This document describes how distributed responds in the face of these failures and other known bugs.

3.19.1 User code failures

When a function raises an error that error is kept and transmitted to the executor on request. Any attempt to gather that result *or any dependent result* will raise that exception.

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```
>>> def div(a, b):
...    return a / b

>>> x = executor.submit(div, 1, 0)
>>> x.result()
ZeroDivisionError: division by zero

>>> y = executor.submit(add, x, 10)
>>> y.result() # same error as above
ZeroDivisionError: division by zero
```

This does not affect the smooth operation of the scheduler or worker in any way.

3.19.2 Closed Network Connections

If the connection to a remote worker unexpectedly closes and the local process appropriately raises an IOError then the scheduler will reroute all pending computations to other workers.

If the lost worker was the only worker to hold vital results necessary for future computations then those results will be recomputed by surviving workers. The scheduler maintains a full history of how each result was produced and so is able to reproduce those same computations on other workers.

This has some fail cases.

- 1. If results depend on impure functions then you may get a different (although still entirely accurate) result
- 2. If the worker failed due to a bad function, for example a function that causes a segmentation fault, then that bad function will repeatedly be called on other workers, and proceed to kill the distributed system, one worker at a time.
- 3. Data "scatter" ed out to the workers is not kept in the scheduler (it is often quite large) and so the loss of this data is irreparable.

3.19.3 Hardware Failures

It is not clear under which circumstances the local process will know that the remote worker has closed the connection. If the socket does not close cleanly then the system will wait for a timeout, roughly three seconds, before marking the worker as failed and resuming smooth operation.

3.19.4 Scheduler Failure

The process containing the scheduler might die. There is currently no persistence mechanism to record and recover the scheduler state. The data will remain on the cluster until cleared.

3.19.5 Restart and Nanny Processes

The executor provides a mechanism to restart all of the workers in the cluster. This is convenient if, during the course of experimentation, you find your workers in an inconvenient state that makes them unresponsive. The Executor.restart method does the following process:

- 1. Sends a soft shutdown signal to all of the coroutines watching workers
- 2. Sends a hard kill signal to each worker's Nanny process, which oversees that worker. This Nanny process terminates the worker process ungracefully and unregisters that worker from the Scheduler.

- 3. Clears out all scheduler state and sets all Future's status to 'cancelled'
- 4. Sends a restart signal to all Nanny processes, which in turn restart clean Worker processes and register these workers with the Scheduler. New workers may not have the same port as their previous iterations. The .nannies dictionary on the Executor serves as an accurate set of aliases if necessary.
- 5. Restarts the scheduler, with clean and empty state

This effectively removes all data and clears out all computations from the scheduler. Any data or computations not saved to persistent storage are lost. This process is very robust to a number of failure modes, including non-responsive or swamped workers but not including full hardware failures.

Currently the user may experience a few error logging messages from Tornado upon closing their session. These can safely be ignored.

3.20 Journey of a Task

We follow a single task through the user interface, scheduler, worker nodes, and back. Hopefully this helps to illustrate the inner workings of the system.

3.20.1 User code

A user computes the addition of two variables already on the cluster, then pulls the result back to the local process.

```
e = Executor('host:port')
x = e.submit(...)
y = e.submit(...)
z = e.submit(add, x, y) # we follow z
print(z.result())
```

3.20.2 Step 1: Executor

z begins its life when the Executor. submit function sends the following message to the Scheduler:

```
{'op': 'update-graph',
  'tasks': {'z': (add, x, y)},
  'keys': ['z']}
```

The executor then creates a Future object with the key 'z' and returns that object back to the user. This happens even before the message has been received by the scheduler. The status of the future says 'pending'.

3.20.3 Step 2: Arrive in the Scheduler

A few milliseconds later, the scheduler receives this message on an open socket.

The scheduler updates its state with this little graph that shows how to compute z.:

```
scheduler.tasks.update[msg['tasks']]
```

The scheduler also updates a lot of other state. Notably, it has to identify that x and y are themselves variables, and connect all of those dependencies. This is a long and detail oriented process that involves updating roughly 10 sets and dictionaries. Interested readers should investigate distributed/scheduler.py::update_state(). While this is fairly complex and tedious to describe rest assured that it all happens in constant time and in about a millisecond.

3.20.4 Step 3: Select a Worker

Once the latter of x and y finishes, the scheduler notices that all of z's dependencies are in memory and that z itself may now run. Which worker should z select? We consider a sequence of criteria:

- 1. First, we quickly downselect to only those workers that have either x or y in local memory.
- 2. Then, we select the worker that would have to gather the least number of bytes in order to get both x and y locally. E.g. if two different workers have x and y and if y takes up more bytes than x then we select the machine that holds y so that we don't have to communicate as much.
- 3. If there are multiple workers that require the minimum number of communication bytes then we select the worker that is the least busy

z considers the workers and chooses one based on the above criteria. In the common case the choice is pretty obvious after step 1. z waits on a stack associated with the chosen worker. The worker may still be busy though, so z may wait a while.

Note: This policy is under flux and this part of this document is quite possibly out of date.

3.20.5 Step 4: Transmit to the Worker

Eventually the worker finishes a task, has a spare core, and z finds itself at the top of the stack (note, that this may be some time after the last section if other tasks placed themselves on top of the worker's stack in the meantime.)

We place z into a worker_queue associated with that worker and a worker_core coroutine pulls it out. z's function, the keys associated to its arguments, and the locations of workers that hold those keys are packed up into a message that looks like this:

This message is serialized and sent across a TCP socket to the worker.

3.20.6 Step 5: Execute on the Worker

The worker unpacks the message, and notices that it needs to have both x and y. If the worker does not already have both of these then it gathers them from the workers listed in the who_has dictionary also in the message. For each key that it doesn't have it selects a valid worker from who_has at random and gathers data from it.

After this exchange, the worker has both the value for x and the value for y. So it launches the computation add (x, y) in a local ThreadPoolExecutor and waits on the result.

In the mean time the worker repeats this process concurrently for other tasks. Nothing blocks.

Eventually the computation completes. The Worker stores this result in its local memory:

```
data['x'] = ...
```

And transmits back a success, and the number of bytes of the result:

```
Worker: Hey Scheduler, 'z' worked great.
I'm holding onto it.
It takes up 64 bytes.
```

The worker does not transmit back the actual value for z.

3.20.7 Step 6: Scheduler Aftermath

The scheduler receives this message and does a few things:

- 1. It notes that the worker has a free core, and sends up another task if available
- 2. If x or y are no longer needed then it sends a message out to relevant workers to delete them from local memory.
- 3. It sends a message to all of the clients that z is ready and so all client Future objects that are currently waiting should, wake up. In particular, this wakes up the z.result() command executed by the user originally.

3.20.8 Step 7: Gather

When the user calls z.result() they wait both on the completion of the computation and for the computation to be sent back over the wire to the local process. Usually this isn't necessary, usually you don't want to move data back to the local process but instead want to keep in on the cluster.

But perhaps the user really wanted to actually know this value, so they called z.result().

The scheduler checks who has z and sends them a message asking for the result. This message doesn't wait in a queue or for other jobs to complete, it starts instantly. The value gets serialized, sent over TCP, and then deserialized and returned to the user (passing through a queue or two on the way.)

3.20.9 Step 8: Garbage Collection

The user leaves this part of their code and the local variable z goes out of scope. The Python garbage collector cleans it up. This triggers a decremented reference on the executor (we didn't mention this, but when we created the Future we also started a reference count.) If this is the only instance of a Future pointing to z then we send a message up to the scheduler that it is OK to release z. The user no longer requires it to persist.

The scheduler receives this message and, if there are no computations that might depend on z in the immediate future, it removes elements of this key from local scheduler state and adds the key to a list of keys to be deleted periodically. Every 500 ms a message goes out to relevant workers telling them which keys they can delete from their local memory. The graph/recipe to create the result of z persists in the scheduler for all time.

3.20.10 Overhead

The user experiences this in about 10 milliseconds, depending on network latency.

3.21 Protocol

The scheduler, workers, and clients pass messages between each other. Semantically these messages encode commands, status updates, and data, like the following:

- Please compute the function sum on the data x and store in y
- The computation y has been completed
- Be advised that a new worker named alice is available for use
- Here is the data for the keys 'x', and 'y'

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In practice we represent these messages with dictionaries/mappings:

```
{'op': 'compute',
  'function': ...
  'args': ['x']}

{'op': 'task-complete',
  'key': 'y',
  'nbytes': 26}

{'op': 'register-worker',
  'address': '192.168.1.42',
  'name': 'alice',
  'ncores': 4}

{'x': b'...',
  'y': b'...'}
```

When we communicate these messages between nodes we need to serialize these messages down to a string of bytes that can then be describilized on the other end to their in-memory dictionary form. For simple cases several options exist like JSON, MsgPack, Protobuffers, and Thrift. The situation is made more complex by concerns like serializing Python functions and Python objects, optional compression, cross-language support, large messages, and efficiency.

This document describes the protocol used by dask.distributed today. Be advised that this protocol changes rapidly as we continue to optimize for performance.

3.21.1 Overview

We may split a single message into multiple message-part to suit different protocols. Generally small bits of data are encoded with MsgPack while large bytestrings are handled specially by a custom format. Each message-part gets its own header, which is always encoded as msgpack. After serializing all message parts we have a sequence of bytestrings or *frames* which we send along the wire, prepended with length information.

The application doesn't know any of this, it just sends us Python dictionaries with various datatypes and we produce a list of bytestrings that get written to a socket. This format is fast both for many frequent messages and for large messages.

3.21.2 MsgPack for Messages

Most messages are encoded with MsgPack, a self describing semi-structured serialization format that is very similar to JSON, but smaller, faster, not human-readable, and supporting of bytestrings and (soon) timestamps. We chose MsgPack as a base serialization format for the following reasons:

- It does not require separate headers, and so is easy and flexible to use which is particularly important in an early stage project like dask.distributed
- It is very fast, much faster than JSON, and there are nicely optimized implementations, particularly within the pandas.msgpack module. With few exceptions (described later) MsgPack does not come anywhere near being a bottleneck, even under heavy use.
- Unlike JSON it supports bytestrings
- It covers the standard set of types necessary to encode most information
- It is widely implemented in a number of languages (see cross language section below)

However, MsgPack fails (correctly) in the following ways:

- It does not provide any way for us to encode Python functions or user defined data types
- It does not support bytestrings greater than 4GB and is generally inefficient for very large messages.

Because of these failings we supplement it with a language-specific protocol and a special case for large bytestrings.

3.21.3 CloudPickle for Functions and Data

Pickle and CloudPickle are Python libraries to serialize almost any Python object, including functions. We use these libraries to transform the users' functions and data into bytes before we include them in the dictionary/map that we pass off to msgpack. In the introductory example you may have noticed that we skipped providing an example for the function argument:

```
{'op': 'compute',
  'function': ...
  'args': ['x']}
```

That is because this value . . . will actually be the result of calling cloudpickle.dumps (myfunction). Those bytes will then be included in the dictionary that we send off to msgpack, which will only have to deal with bytes rather than obscure Python functions.

3.21.4 Cross Language Specialization

The Client and Workers must agree on a language-specific serialization format. In the standard dask.distributed client and worker objects this ends up being the following:

```
bytes = cloudpickle.dumps(obj, protocol=pickle.HIGHEST_PROTOCOL)
obj = cloudpickle.loads(bytes)
```

This varies between Python 2 and 3 and so your client and workers must match their Python versions and software environments.

However, the Scheduler never uses the language-specific serialization and instead only deals with MsgPack. If the client sends a pickled function up to the scheduler the scheduler will not unpack function but will instead keep it as bytes. Eventually those bytes will be sent to a worker, which will then unpack the bytes into a proper Python function. Because the Scheduler never unpacks language-specific serialized bytes it may be in a different language.

The client and workers must share the same language and software environment, the scheduler may differ.

This has a few advantages:

- 1. The Scheduler is protected from unpickling unsafe code
- 2. The Scheduler can be run under pypy for improved performance. This is only useful for larger clusters.
- 3. We could conceivably implement workers and clients for other languages (like R or Julia) and reuse the Python scheduler. The worker and client code is fairly simple and much easier to reimplement than the scheduler, which is complex.
- 4. The scheduler might some day be rewritten in more heavily optimized C or Go

3.21.5 Compression

Fast compression libraries like LZ4 or Snappy may increase effective bandwidth by compressing data before sending and decompressing it after reception. This is especially valuable on lower-bandwidth networks.

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If either of these libraries is available (we prefer LZ4 to Snappy) then for every message greater than 1kB we try to compress the message and, if the compression is at least a 10% improvement, we send the compressed bytes rather than the original payload. We record the compression used within the header as a string like 'lz4' or 'snappy'.

To avoid compressing large amounts of uncompressable data we first try to compress a sample. We take 10kB chunks from five locations in the dataset, arrange them together, and try compressing the result. If this doesn't result in significant compression then we don't try to compress the full result.

3.21.6 Header

The header is a small dictionary encoded in msgpack that includes some metadata about the message, such as compression.

3.21.7 Large Bytestrings

Whenever a message comes in with very large byte values like the following:

```
{'key': 'x',
  'address': 'alice',
  'data-1': b'...' # very long bytestring
  'data-2': b'...' # very long bytestring
}
```

We separate the message into two messages, one encoding all of the large bytestrings, and one encoding everything else:

```
{'key': 'x', 'addresss': 'alice'}
{'data-1': b'...', 'data-2': b'...'}
```

The first message we pass normally with msgpack, the second we pass in multiple parts, including a header that contains the keys and compression used for each value:

```
{'keys': ['data-1', 'data-2'],
  'compression': ['lz4', None]}
b'...'
b'...'
```

3.21.8 Frames

At the end of the pipeline we have a sequence of bytestrings or frames. We need to tell the receiving end how many frames there are and how long each these frames are. We order the frames and lengths of frames as follows:

- 1. The number of frames, stored as an 8 byte unsigned integer
- 2. The length of each frame, each stored as an 8 byte unsigned integer
- 3. Each of the frames

In the following sections we describe how we create these frames.

3.21.9 Performance

For large numpy arrays we currently suffer three memory copies. On a nice machine this ends up being a 1-1.5 GB/s bottleneck, which is almost always faster than the network bandwidth. These copies come from NumPy (two memcopies) and Tornado (one memcopy).

For small messages we generally serialize in around 5 microseconds.

3.22 Work Stealing

Some tasks prefer to run on certain workers. This may be because that worker holds data dependencies of the task or because the user has expressed a loose desire that the task run in a particular place. Occasionally this results in a few very busy workers and several idle workers. In this situation the idle workers may choose to steal work from busy workers, even if stealing work requires the costly movement of data.

This is a performance optimization and not required for correctness. Work stealing provides robustness in many ad-hoc cases, but can also backfire when we steal the wrong tasks and reduce performance.

3.22.1 Task criteria for stealing

If a task has been specifically restricted to run on particular workers (such as is the case when special hardware is required) then we do not steal. Barring this case, stealing usually occurs for tasks that have been assigned to a particular worker because that worker holds the data necessary to compute the task.

Stealing is profitable when the computation time for a task is much longer than the communication time of the task's dependencies. It is also good long term if stealing causes highly-sought-after data to be replicated on more workers.

Bad example

We do not want to steal tasks that require moving a large dependent piece of data across a wire from the victim to the thief if the computation is fast. We end up spending far more time in communication than just waiting a bit longer and giving up on parallelism.

```
[data] = e.scatter([np.arange(1000000000)])
x = e.submit(np.sum, data)
```

Good example

We do want to steal task tasks that only need to move dependent pieces of data, especially when the computation time is expensive (here 100 seconds.)

```
[data] = e.scatter([100])
x = e.submit(sleep, data)
```

Fortunately we often know both the number of bytes of dependencies (as reported by calling sys.getsizeof on the workers) and the runtime cost of previously seen functions.

3.22.2 When do we worksteal

The scheduler maintains a set of idle workers and a set of saturated workers. At various events, such as when new tasks arrive from the client, when new workers arrive, or when we learn that workers have completed a set of tasks, we play these two sets of idle and saturated workers against each other.

3.22.3 Choosing tasks to steal

Occupied workers maintain a stack of excess work. The tasks at the top of this stack are prioritized to be run by that worker before the tasks at the bottom.

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Ideally we choose the worker with the *largest* stack of excess work and then select the task at the *bottom* of this stack, hopefully starting a new sequence of computations that are somewhat unrelated to what the busy worker is currently working on.

All operations in the scheduler endeavor to be computed in constant time (or linear time relative to the number of processed tasks.) We can pull from the bottom of the stack in constant time by implementing each worker's stack as a collections.deque. However, we currently do not maintain the data structures necessary to efficiently find the most occupied workers. Common solutions, like maintaining priority queue of workers by stack length add a log (n) cost to the common case.

Instead we just call next (iter(saturated_workers)) and allow Python to iterate through the set of saturated workers however it prefers.

3.23 Scheduler Plugins

class distributed.diagnostics.plugin.SchedulerPlugin

Interface to extend the Scheduler

The scheduler operates by triggering and responding to events like task_finished, update_graph, task erred, etc..

A plugin enables custom code to run at each of those same events. The scheduler will run the analagous methods on this class when each event is triggered. This runs user code within the scheduler thread that can perform arbitrary operations in synchrony with the scheduler itself.

Plugins are often used for diagnostics and measurement, but have full access to the scheduler and could in principle affect core scheduling.

To implement a plugin implement some of the methods of this class and add the plugin to the scheduler with Scheduler.add_plugin(myplugin).

Examples

```
>>> class Counter(SchedulerPlugin):
...     def __init__(self):
...          self.counter = 0
...
...     def task_finished(self, scheduler, key, worker, nbytes):
...          self.counter += 1
...
...     def restart(self, scheduler):
...          self.counter = 0
```

```
>>> c = Counter()
>>> scheduler.add_plugin(c)
```

```
restart (scheduler, **kwargs)
```

Run when the scheduler restarts itself

```
task_erred (scheduler, key=None, worker=None, exception=None, **kwargs)
Run when a task is reported failed
```

```
task_finished (scheduler, key=None, worker=None, nbytes=None, **kwargs)
Run when a task is reported complete
```

```
update_graph (scheduler, dsk=None, keys=None, restrictions=None, **kwargs)
Run when a new graph / tasks enter the scheduler
```

3.24 Related Work

Writing the "related work" for a project called "distributed", is a Sisyphean task. We'll list a few notable projects that you've probably already heard of down below.

You may also find the dask comparison with spark of interest.

3.24.1 Big Data World

- The venerable Hadoop provides batch processing with the MapReduce programming paradigm. Python users typically use Hadoop Streaming or MRJob.
- Spark builds on top of HDFS systems with a nicer API and in-memory processing. Python users typically use PySpark.
- Storm provides streaming computation. Python users typically use streamparse.

This is a woefully inadequate representation of the excellent work blossoming in this space. A variety of projects have come into this space and rival or complement the projects above. Still, most "Big Data" processing hype probably centers around the three projects above, or their derivatives.

3.24.2 Python Projects

There are dozens of Python projects for distributed computing. Here we list a few of the more prominent projects that we see in active use today.

Task scheduling

- Celery: An asynchronous task scheduler, focusing on real-time processing.
- Luigi: A bulk big-data/batch task scheduler, with hooks to a variety of interesting data sources.

Ad hoc computation

- IPython Parallel: Allows for stateful remote control of several running ipython sessions.
- Scoop: Implements the concurrent.futures API on distributed workers. Notably allows tasks to spawn more tasks.

Direct Communication

- MPI4Py: Wraps the Message Passing Interface popular in high performance computing.
- PyZMQ: Wraps ZeroMQ, the gentleman's socket.

Venerable

There are a couple of older projects that often get mentioned

- Dispy: Embarrassingly parallel function evaluation
- Pyro: Remote objects / RPC

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3.24.3 Relationship

In relation to these projects distributed...

- Supports data-local computation like Hadoop and Spark
- Uses a task graph with data dependencies abstraction like Luigi
- In support of ad-hoc applications, like IPython Parallel and Scoop

3.24.4 In depth comparison to particular projects

IPython Parallel

Short Description

IPython Parallel is a distributed computing framework from the IPython project. It uses a centralized hub to farm out jobs to several ipengine processes running on remote workers. It communicates over ZeroMQ sockets and centralizes communication through the central hub.

IPython parallel has been around for a while and, while not particularly fancy, is quite stable and robust.

IPython Parallel offers parallel map and remote apply functions that route computations to remote workers

```
>>> view = Client(...)[:]
>>> results = view.map(func, sequence)
>>> result = view.apply(func, *args, **kwargs)
>>> future = view.apply_async(func, *args, **kwargs)
```

It also provides direct execution of code in the remote process and collection of data from the remote namespace.

```
>>> view.execute('x = 1 + 2')
>>> view['x']
[3, 3, 3, 3, 3, 3]
```

Brief Comparison

Distributed and IPython Parallel are similar in that they provide map and apply/submit abstractions over distributed worker processes running Python. Both manage the remote namespaces of those worker processes.

They are dissimilar in terms of their maturity, how worker nodes communicate to each other, and in the complexity of algorithms that they enable.

Distributed Advantages

The primary advantages of distributed over IPython Parallel include

- 1. Peer-to-peer communication between workers
- 2. Dynamic task scheduling

Distributed workers share data in a peer-to-peer fashion, without having to send intermediate results through a central bottleneck. This allows distributed to be more effective for more complex algorithms and to manage larger datasets in a more natural manner. IPython parallel does not provide a mechanism for workers to communicate with each other, except by using the central node as an intermediary for data transfer or by relying on some other medium, like a shared file system. Data transfer through the central node can easily become a bottleneck and so IPython parallel has been mostly helpful in embarrassingly parallel work (the bulk of applications) but has not been used extensively for more sophisticated algorithms that require non-trivial communication patterns.

The distributed executor includes a dynamic task scheduler capable of managing deep data dependencies between tasks. The IPython parallel docs include a recipe for executing task graphs with data dependencies. This

same idea is core to all of distributed, which uses a dynamic task scheduler for all operations. Notably, distributed. Future objects can be used within submit/map/get calls before they have completed.

```
>>> x = executor.submit(f, 1) # returns a future
>>> y = executor.submit(f, 2) # returns a future
>>> z = executor.submit(add, x, y) # consumes futures
```

The ability to use futures cheaply within submit and map methods enables the construction of very sophisticated data pipelines with simple code. Additionally, distributed can serve as a full dask task scheduler, enabling support for distributed arrays, dataframes, machine learning pipelines, and any other application build on dask graphs. The dynamic task schedulers within distributed are adapted from the dask task schedulers and so are fairly sophisticated/efficient.

IPython Parallel Advantages

IPython Parallel has the following advantages over distributed

- 1. Maturity: IPython Parallel has been around for a while.
- 2. Explicit control over the worker processes: IPython parallel allows you to execute arbitrary statements on the workers, allowing it to serve in system administration tasks.
- 3. Deployment help: IPython Parallel has mechanisms built-in to aid deployment on SGE, MPI, etc.. Distributed does not have any such sugar, though is fairly simple to set up by hand.
- 4. Various other advantages: Over the years IPython parallel has accrued a variety of helpful features like IPython interaction magics, @parallel decorators, etc..

concurrent.futures

The distributed. Executor API is modeled after concurrent. futures and PEP-3184. It has a few notable differences:

- distributed accepts Future objects within calls to submit/map. It is preferable to submit Future objects directly rather than wait on them before submission.
- The map function returns Future objects, not concrete results. The map function returns immediately.
- It is not yet possible to cancel a Future (though this is theoretically possible please raise an issue if this is of concrete importance to you.)
- Distributed generally does not support timeouts or callbacks

distributed. Compatible Executor is a subclass of distributed. Executor that does conform to the concurrent. futures API, allowing it to be used as a drop-in replacement for other Executors using the common API.

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