

# UMD DATA605 - Big Data Systems

## Python Dask

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**v1.1**

# Dask: Resources

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- Web resources:
  - Dask project
  - Dask examples
- Tutorial
  - Dask\_tutorial
  - Dask\_advanced\_tutorial
- Class project
- Mastery
  - Data science with Python and Dask, 2019
    - Amazon

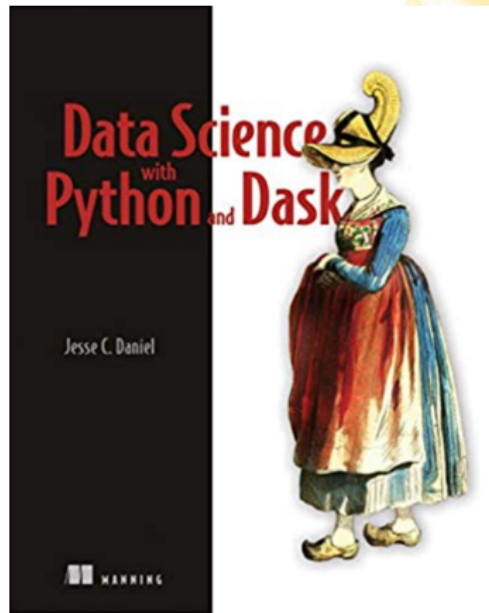


Figure 1: alt\_text

# Dataset Size Issues

- **Small datasets (< 1 GB)**
  - Fits into RAM
  - Manipulation doesn't require paging to disk
- **Medium dataset (< 1TB)**
  - Doesn't fit into RAM
  - Fits into local disk
    - Performance penalty imposed by using local disk
  - Need multiple CPU cores
    - Difficult to take advantage of parallelism with Python / Pandas
- **Large dataset (> 1TB)**
  - Doesn't fit into RAM
  - Doesn't fit into local disk



# Dataset Size Issues

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- **Small datasets**
  - < 1 GB
- **Medium dataset**
  - < 1TB
- **Large dataset**
  - 1TB
- **The thresholds are fuzzy and changing over time**
  - E.g., you can scale the computer 10x and get 10x bigger data sets
- **Problem with scaling datasets**
  - Long run times
  - Rewriting code in different language / API for datasets of different size
  - Need to think about *what to do it* and *how to do it* efficiently
  - Cumbersome framework (Pandas easy, Hadoop difficult)

# Dask

- **Dask is written in Python**

- It scales natively Numpy, Pandas, sklearn
- Dask objects are wrappers (don't just mirror the interface) objects from the respective libraries (e.g., Pandas DataFrame, numpy array)
- Parallel parts are called "chunks" or "partitions"
  - Are queued to be worked on
  - Shipped between machines
  - Worked locally on a machine

- **Pros**

- Users don't need to learn a new language, but can use familiar interfaces
- Can focus on writing code that is optimized for parallelism
  - Dask does the heavy lifting

- **Scaling Dask is easy**

- Users can write a prototype task on a local machines and use a cluster when needed



Figure 2: alt\_text



SCIENCE  
ACADEMY

No need to refactor existing code

# Dask Layers

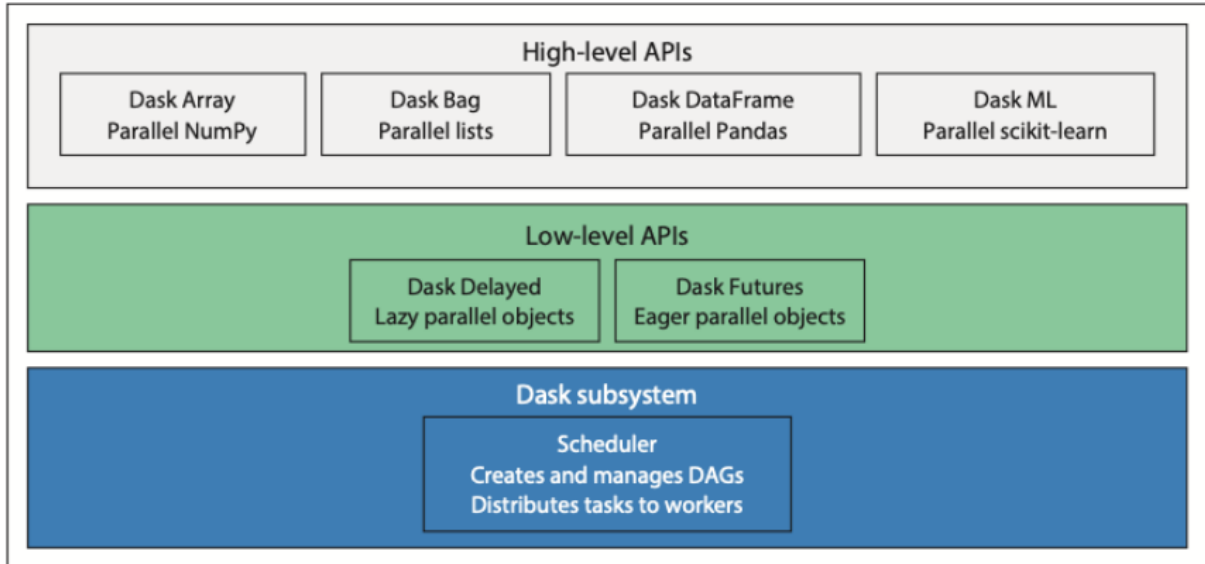


Figure 3: alt\_text

# Scaling Up vs Scaling Out

## • Scaling up

- = replace equipment with larger, faster equipments
  - E.g., buy a larger pot, replace knife with food processor

### • Pros

- You got better hardware, nothing else needs to change (e.g., code)

### • Cons

- There will be a time where you exceed the capacity of the current machines
- Cost: more powerful machines are expensive



## • Scaling out

- = divide the work between many workers in parallel
  - E.g., buy more pots and hire more cooks

### • Pros

- Task scheduler organizes computation, assigning workers to each task
- More cost-effective solution since no specialized hardware is needed



### • Cons

- Need to write code to expose parallelism
- Costs of maintaining a cluster



# Dask: Computation

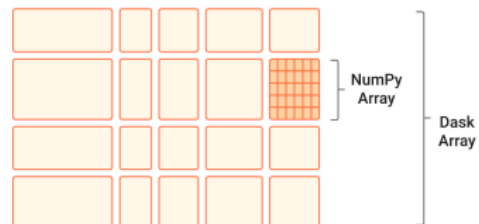
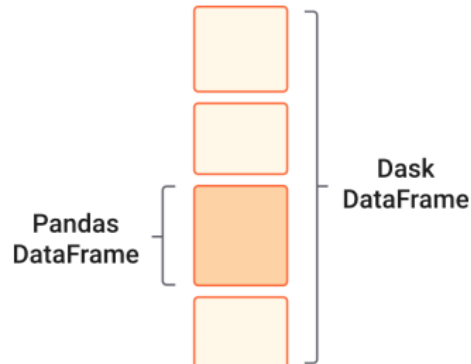
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- **Lazy computations**
  - User defines the transformations on the data
  - No need to wait for one computation to finish before defining the next
  - Avoid loading the entire data in memory by operating in chunks
  - E.g.,
    - Split a 2GB file into 32 64MB chunks
    - Operate on 8 chunks at a time on each server
    - The max memory consumption doesn't exceed  $512\text{MB} = (8 \times 32)$
  - Each task tracks object dimensions and data types
    - No code is executed
- **compute()**
  - Running a computation (aka materializing) `''' python missing_count_pct = missing_count.compute()'''`
- **persist()**
  - As soon as a node in the graph emits results, its intermediate work is discarded to minimize memory usage
  - If we need to do additional computation on intermediate nodes we need to re-run the graph
  - **persist()** tells Dask to keep the intermediate result in memory
  - This speeds up a large and complex DAG that needs to be reused many



# Dask: Data Structures

- **Dask DataFrame**  
implements Pandas DataFrame
  - Tabular / relational data
- **Dask Array** implements numpy ndarray
  - Multidimensional array\*\*
- **Dag Bag** coordinates Python lists of objects
  - Parallelize computations on unstructured or semi-structured data



```
[1, 2, 3, 4, 5]  
[1, 2, 3] [4, 5]
```

## Dask Reading Data

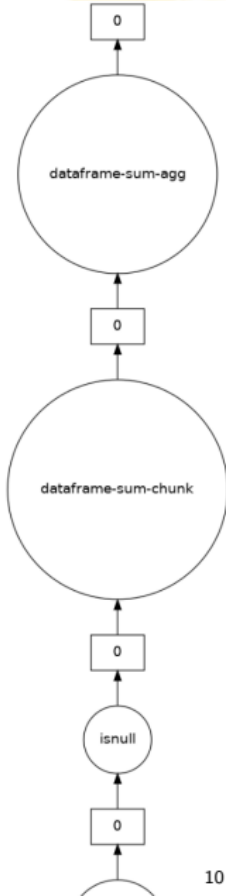
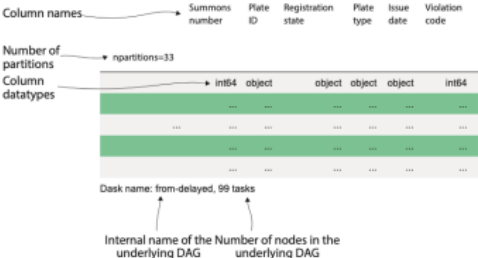
- 1

## dask.dataframe.read\_csv()

- Doesn't load the data in memory with
  - Tries to infer the types of the columns
    - By randomly sampling some data
    - Best to set the data types
    - Even better is to use Parquet since it stores data and types together

- Partitions = chunks of data that can be worked independently

- E.g., 33 partitions
  - Graph is composed of 99 tasks
  - Each partition reads ENCE data, splits data, ACADEMY initializes df object



# Low Level APIs: Delayed

- Handle computations that don't fit in native Dask data structures (e.g., Dask DataFrame)
- In the example below there is parallelism that can be exploited

```
def inc(x):
    return x + 1

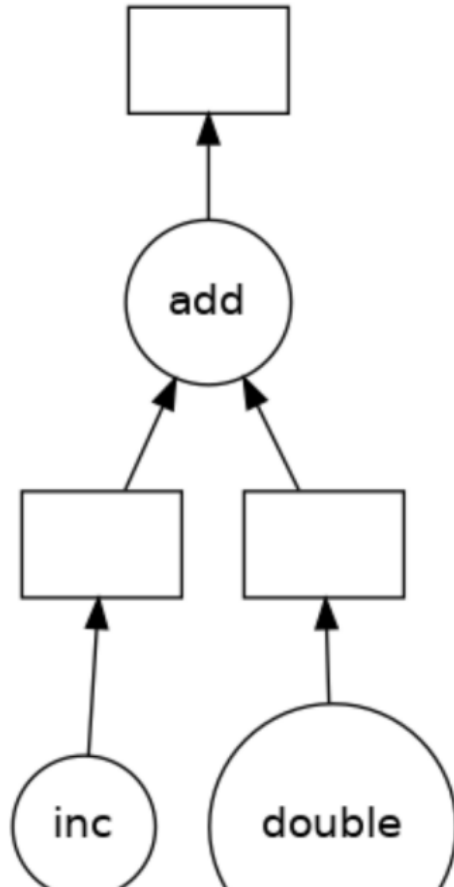
def double(x):
    return x * 2

def add(x, y):
    return x + y

data = [1, 2, 3, 4, 5]

output = []
for x in data:
    #  $(x + 1) + (x * 2) = 3x + 1$ 
    a = inc(x)
    b = double(x)
    c = add(a, b)
    # 1 -> 4
    # 2 -> 7
    # 3 -> 10
    # 4 -> 13
    # 5 -> 16
    output.append(c)

# 4 + 7 + 10 + 13 + 16 = 20 + 20 + 10 = 50
total = sum(output)
print(total)
```

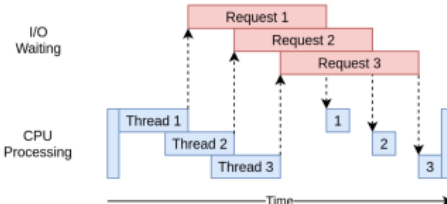


# Low Level APIs: Futures

- In parallel programming, a “future” encapsulates the asynchronous execution of a callable, representing the eventual result of the operation
- Futures is the most general way of specifying concurrency in Dask
  - Everything can be expressed in terms of futures
  - User can specify what's blocking and what's not blocking
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## Python **concurrent.futures**

- High-level interface for asynchronously executing callables
- Thread pool or Process



```
def inc(x):  
    return x + 1
```

```
def add(x, y):  
    return x + y
```

```
a = client.submit(inc, 10)  
b = client.submit(inc, 20)
```

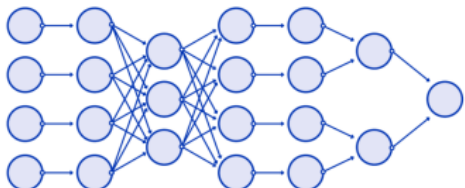
```
>>> a  
<Future: status: pending, key: inc-b8aaef26b99466a7a1980efafade6701d>  
>>> a  
<Future: status: finished, type: int, key: inc-b8aaef26b99466a7a1980efafade6701d>  
>>> a.result() # blocks until task completes and data arrives  
11
```

# Different Types of Parallel Workload

- Break program in medium-size tasks of computation
  - E.g., a function call

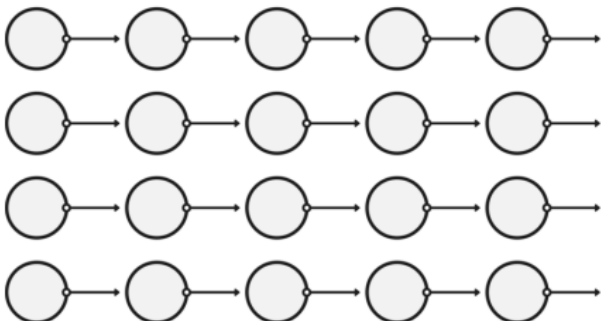
MapReduce

Hadoop/Spark/Dask



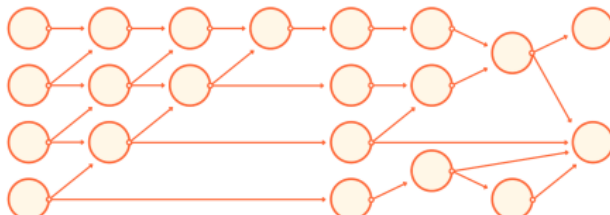
## Embarrassingly Parallel

Hadoop/Spark/Dask/Airflow/Prefect



## Full Task Scheduling

Dask/Airflow/Prefect



# Encoding Task Graph

- Dask encodes tasks in terms of Python dicts and functions :: columns ::::

```
def inc(i):  
    return i + 1
```

```
def add(a, b):  
    return a + b
```

```
x = 1  
y = inc(x)  
z = add(y, 10)
```

# Task Scheduling

- Data collections (Bags, Arrays, DataFrame) and their operations create task graphs
  - Nodes in the task graph are Python functions
  - Edges are dependencies (e.g., output from one task used as input in another task)
- Task graphs are scheduled for execution
- Single-machine scheduler
  - Use local process or thread pool
  - Simple but it can only run on a single machine
- Distributed scheduler
  - It can run locally or distributed across a cluster

Collections

(create task graphs)

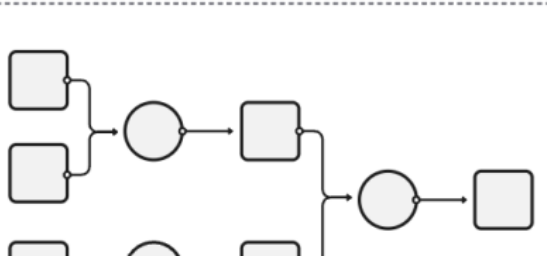


Task Graph



Schedulers

(execute task graphs)



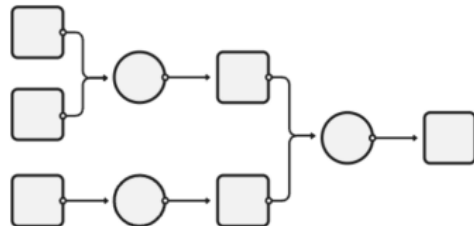
Single-machine  
(threads, processes,  
synchronous)

Distributed



# Task Scheduling

- Dask task scheduler orchestrates the work dynamically
  - Not a static scheduling of operations like a relational DB
  - When the computation takes place, Dask dynamically assesses:
    - What tasks has been completed
    - What tasks is left to do
    - What resources (CPUs) are free
    - Where the data is located



- This dynamic approach handles a variety issues:

- Worker failure
  - Just re-run
- Workers completing work at different speeds

# Dask vs Spark

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- Spark has
- **Pros**
  - Popular framework for analyzing large datasets
  - In-memory alternative to MapReduce / Hadoop
- **Cons**
  - Spark is a Java library, supporting Python through PySpark API
    - Python code is executed on JVM through py4j
    - Difficult to debug since execution occurs outside Python
  - Different DataFrame API than Pandas
    - Learn how to do things “the Spark way”
    - You might need to implement things twice to go from exploratory analysis to large experiments / production
  - Optimized for MapReduce operations over a collection
  - Difficult to set-up and configure

# Tutorial

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Tutorial - From the official documentation

<https://docs.dask.org/en/stable/10-minutes-to-dask.html>