

# Numerical optimization (cont.)

## Lecture 13

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```
1 .reveal code {  
2   white-space: pre;  
3 }
```

# Method Summary

SciPy Method	Description	Gradient	Hessian
—	Gradient Descent (naive)	✓	✗
—	Newton's method (naive)	✓	✓
—	Conjugate Gradient (naive)	✓	✓
"CG"	Nonlinear Conjugate Gradient (Polak and Ribiere variation)	✓	✗
"Newton-CG"	Truncated Newton method (Newton w/ CG step direction)	✓	Optional
"BFGS"	Broyden, Fletcher, Goldfarb, and Shanno (Quasi-newton method)	Optional	✗
"L-BFGS-B"	Limited-memory BFGS (Quasi-newton method)	Optional	✗
"Nelder-Mead"	Nelder-Mead simplex reflection method	✗	✗

# Methods collection

```
1 def define_methods(x0, f, grad, hess, tol=1e-8):
2     return {
3         "naive_newton": lambda: newtons_method(x0, f, grad, hess, tol=tol),
4         "naive_cg": lambda: conjugate_gradient(x0, f, grad, hess, tol=tol),
5         "CG": lambda: optimize.minimize(f, x0, jac=grad, method="CG", tol=tol),
6         "newton-cg": lambda: optimize.minimize(f, x0, jac=grad, hess=None, method="Newton-CG", tol=tol),
7         "newton-cg w/ H": lambda: optimize.minimize(f, x0, jac=grad, hess=hess, method="Newton-CG", tol=tol),
8         "bfgs": lambda: optimize.minimize(f, x0, jac=grad, method="BFGS", tol=tol),
9         "bfgs w/o G": lambda: optimize.minimize(f, x0, method="BFGS", tol=tol),
10        "l-bfgs-b": lambda: optimize.minimize(f, x0, method="L-BFGS-B", tol=tol),
11        "nelder-mead": lambda: optimize.minimize(f, x0, method="Nelder-Mead", tol=tol)
12    }
```

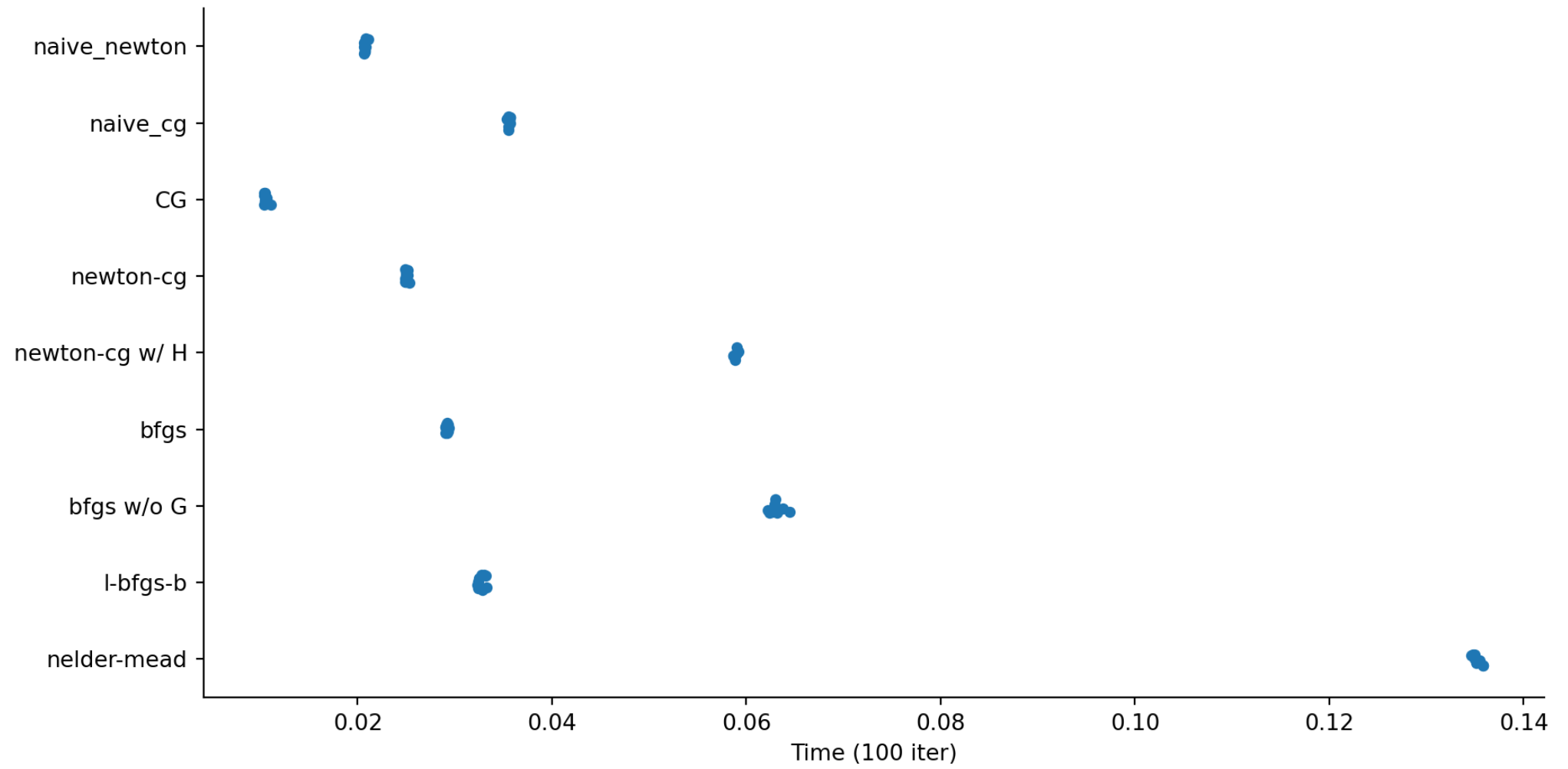
# Method Timings

```
1 x0 = (1.6, 1.1)
2 f, grad, hess = mk_quad(0.7)
3 methods = define_methods(x0, f, grad, hess)
4
5 df = pd.DataFrame({
6     key: timeit.Timer(methods[key]).repeat(10, 100) for key in methods
7 })
8
9 df
```

# Method Timings

	naive_newton	naive_cg	CG	...	bfgs w/o G	l-bfgs-b	nelder-mead
0	0.021096	0.035406	0.011022	...	0.063204	0.032869	0.135478
1	0.020689	0.035320	0.010575	...	0.063190	0.032366	0.134706
2	0.020797	0.035483	0.010438	...	0.062650	0.033036	0.134936
3	0.020783	0.035671	0.010599	...	0.062381	0.032441	0.134595
4	0.020857	0.035638	0.010403	...	0.063007	0.032527	0.135014
5	0.020765	0.035697	0.010402	...	0.064476	0.032307	0.134793
6	0.020727	0.035670	0.010464	...	0.062205	0.033232	0.134989
7	0.020650	0.035522	0.010461	...	0.063753	0.032421	0.135109
8	0.020701	0.035572	0.010389	...	0.063448	0.032707	0.135807
9	0.020657	0.035525	0.010470	...	0.062930	0.033151	0.135847

[10 rows x 9 columns]



# Timings across cost functions

```

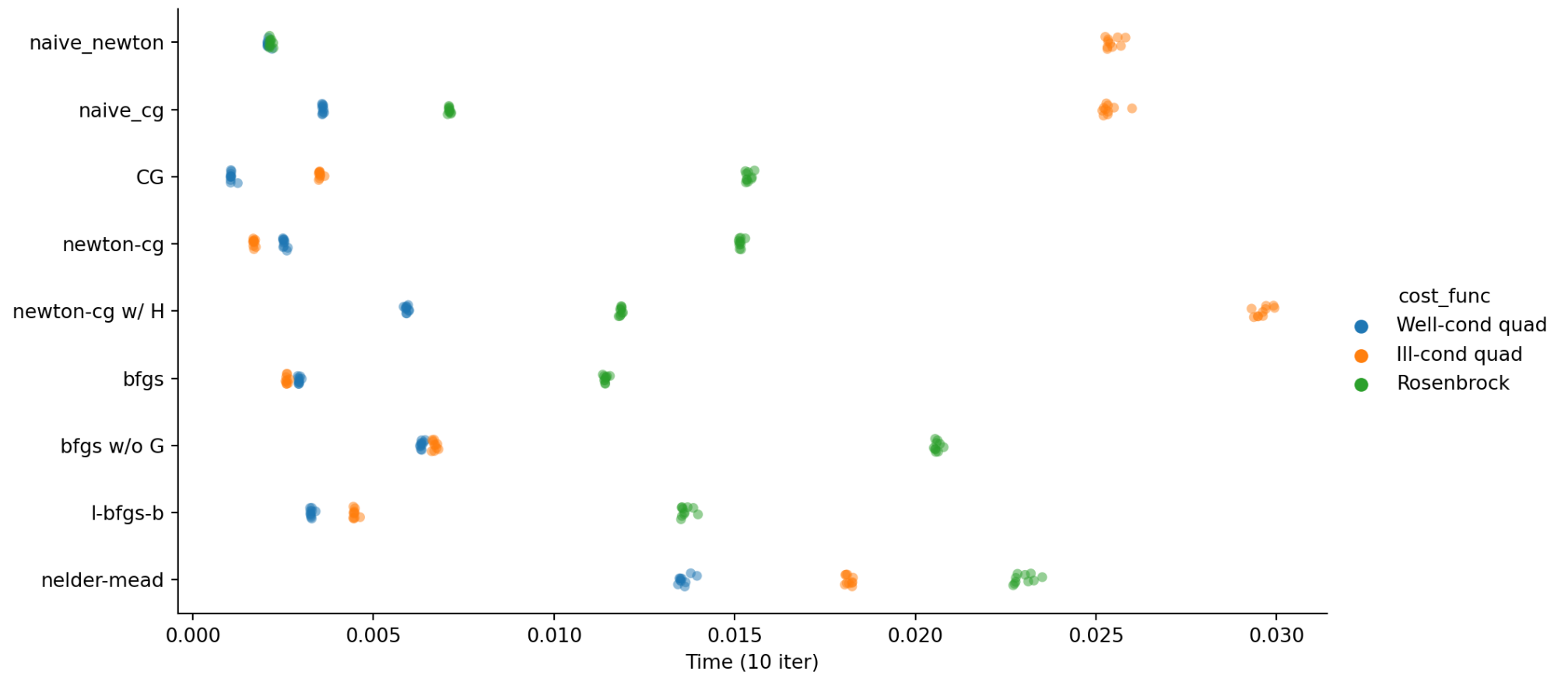
1 def time_cost_func(n, x0, name, cost_func, *args):
2     x0 = (1.6, 1.1)
3     f, grad, hess = cost_func(*args)
4     methods = define_methods(x0, f, grad, hess)
5
6     return ( pd.DataFrame({
7         key: timeit.Timer(
8             methods[key]
9         ).repeat(n, n)
10         for key in methods
11     })
12     .melt()
13     .assign(cost_func = name)
14 )
15
16 df = pd.concat([
17     time_cost_func(10, x0, "Well-cond quad", mk_quad),
18     time_cost_func(10, x0, "Ill-cond quad", mk_quad),
19     time_cost_func(10, x0, "Rosenbrock", mk_rosenb)
20 ])
21
22 df

```

	variable	value	cost_func
0	naive_newton	0.002192	Well-cond quad
1	naive_newton	0.002107	Well-cond quad
2	naive_newton	0.002074	Well-cond quad
3	naive_newton	0.002083	Well-cond quad
4	naive_newton	0.002064	Well-cond quad
..	...	...	...
85	nelder-mead	0.022829	Rosenbrock
86	nelder-mead	0.022748	Rosenbrock
87	nelder-mead	0.022783	Rosenbrock
88	nelder-mead	0.022777	Rosenbrock
89	nelder-mead	0.022710	Rosenbrock

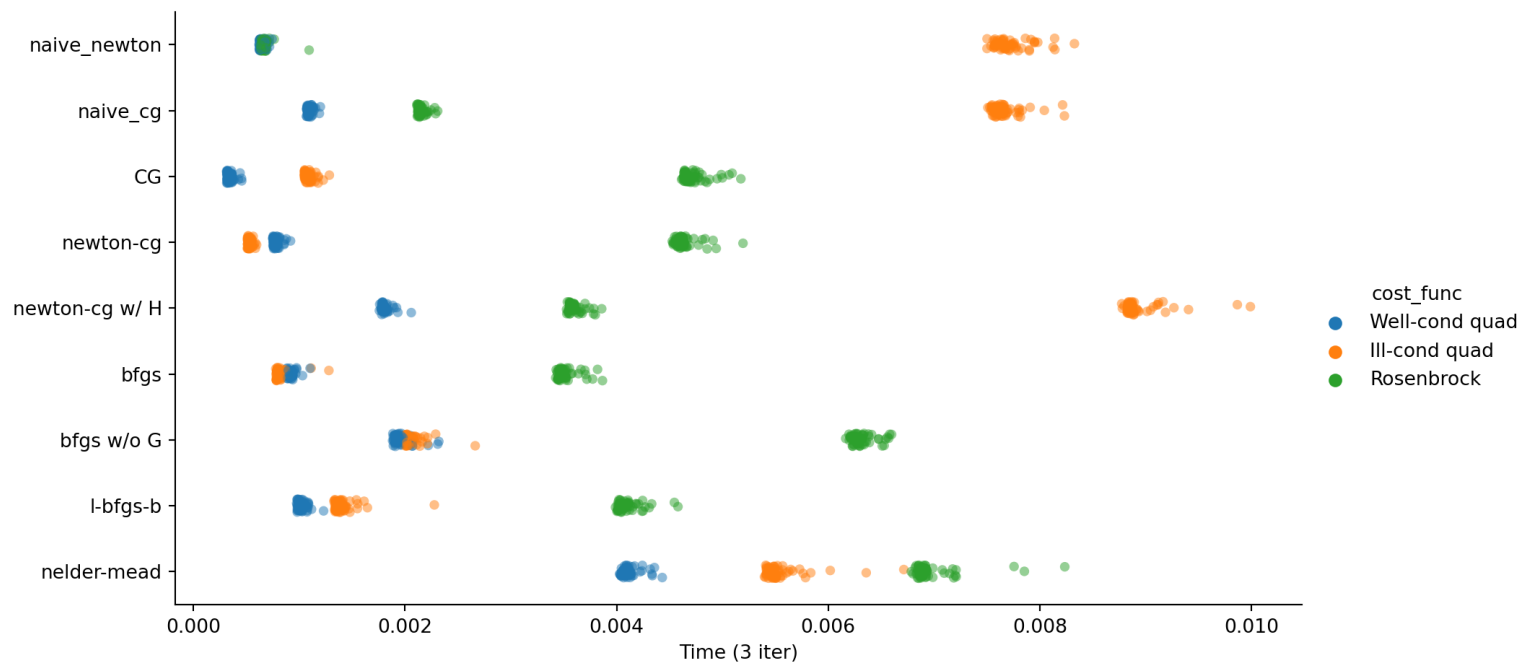
[270 rows x 3 columns]





# Random starting locations

```
1 x0s = np.random.default_rng(seed=1234).uniform(-2,2, (20,2))
2
3 df = pd.concat([
4     pd.concat([
5         time_cost_func(3, x0, "Well-cond quad", mk_quad, 0.7),
6         time_cost_func(3, x0, "Ill-cond quad", mk_quad, 0.02),
7         time_cost_func(3, x0, "Rosenbrock", mk_rosenbrock)
8     ])
9     for x0 in x0s
10 ])
```



# Profiling - BFGS (cProfile)

```
1 import cProfile
2
3 f, grad, hess = mk_quad(0.7)
4 def run():
5     optimize.minimize(fun = f, x0 = (1.6, 1.1), jac=grad, method="BFGS", tol=1e-11)
6
7 cProfile.run('run()', sort="tottime")
```

1293 function calls in 0.001 seconds

Ordered by: internal time

ncalls	tottime	percall	cumtime	percall	filename:lineno(function)
9	0.000	0.000	0.001	0.000	_linesearch.py:31(line_search_wolfel)
1	0.000	0.000	0.001	0.001	_optimize.py:1318(_minimize_bfgs)
58	0.000	0.000	0.000	0.000	{method 'reduce' of 'numpy.ufunc' objects}
152	0.000	0.000	0.000	0.000	{built-in method numpy.core._multiarray_umath.implement_array_
9	0.000	0.000	0.000	0.000	_linesearch.py:91(scalar_search_wolfel)
10	0.000	0.000	0.000	0.000	<string>:2(f)
37	0.000	0.000	0.000	0.000	fromnumeric.py:69(_wrapreduction)
21	0.000	0.000	0.000	0.000	shape_base.py:23(atleast_1d)
26	0.000	0.000	0.000	0.000	_optimize.py:235(vecnorm)
10	0.000	0.000	0.000	0.000	<string>:8(gradient)
20	0.000	0.000	0.000	0.000	numeric.py:2407(array_equal)
1	0.000	0.000	0.001	0.001	_minimize.py:45(minimize)
26	0.000	0.000	0.000	0.000	fromnumeric.py:2188(sum)
1	0.000	0.000	0.000	0.000	_differentiable_functions.py:86(__init__)

51	0.000	0.000	0.000	0.000	<__array_function__ internals>:177(dot)
84	0.000	0.000	0.000	0.000	{built-in method numpy.asarray}
9	0.000	0.000	0.000	0.000	_linesearch.py:77(derphi)
9	0.000	0.000	0.000	0.000	_linesearch.py:73(phi)

# Profiling - BFGS (pyinstrument)

```
1 from pyinstrument import Profiler
2
3 f, grad, hess = mk_quad(0.7)
4
5 profiler = Profiler(interval=0.00001)
6
7 profiler.start()
8 opt = optimize.minimize(fun = f, x0 = (1.6, 1.1), jac=grad, method="BFGS", tol=1e-11)
9 p = profiler.stop()
10
11 profiler.print(show_all=True)
```

```

_      . _   _/_   _ _   _ _/_   Recorded: 10:56:16   Samples:   253
/_//_//_// /_\ / //_// / //_'// //   Duration: 0.004     CPU time: 0.004
/_   _/                                     v4.4.0
```

Program: /opt/homebrew/Cellar/python@3.10/3.10.10\_1/libexec/bin/python3

0.004 MainThread <thread>:8553529664

|- 0.002 [self] None

`- 0.002 <module> <string>:1

  `- 0.002 minimize scipy/optimize/\_minimize.py:45

    `- 0.001 \_minimize\_bfgs scipy/optimize/\_optimize.py:1318

      |- 0.001 \_line\_search\_wolfel2 scipy/optimize/\_optimize.py:1144

        |- 0.001 line\_search\_wolfel1 scipy/optimize/\_linesearch.py:31

          |- 0.001 scalar\_search\_wolfel1 scipy/optimize/\_linesearch.py:91

```

|   |   |- 0.000 phi    scipy/optimize/_linesearch.py:73
|   |   |- 0.000 ScalarFunction.fun    scipy/optimize/_differentiable_functions.py:264
|   |   |   |- 0.000 ScalarFunction._update_fun    scipy/optimize/_differentiable_functions.py:249
|   |   |   |   |- 0.000 update_fun    scipy/optimize/_differentiable_functions.py:154
|   |   |   |   |   |- 0.000 fun_wrapped    scipy/optimize/_differentiable_functions.py:132
|   |   |   |   |   |   |- 0.000 f    <string>:2
|   |   |   |   |   |   |   |- 0.000 sum    <__array_function__ internals>:177
|   |   |   |   |   |   |   |   |- 0.000 sum    numpy/core/fromnumeric.py:2188

```

# Profiling - Nelder-Mead

```
1 from pyinstrument import Profiler
2
3 f, grad, hess = mk_quad(0.7)
4
5 profiler = Profiler(interval=0.00001)
6
7 profiler.start()
8 opt = optimize.minimize(fun = f, x0 = (1.6, 1.1), method="Nelder-Mead", tol=1e-11)
9 p = profiler.stop()
10
11 profiler.print(show_all=True)
```

```

_      . _   _/_   _ _   _ _/_   Recorded: 10:56:16   Samples:  558
/_/_/_/_/_/_/_/_/_/_/_/_/_/_/_   Duration: 0.007      CPU time: 0.007
/_   _/                                     v4.4.0
```

Program: /opt/homebrew/Cellar/python@3.10/3.10.10\_1/libexec/bin/python3

```
0.007 MainThread  <thread>:8553529664
|- 0.005 <module>  <string>:1
|  `-- 0.005 minimize  scipy/optimize/_minimize.py:45
|      `-- 0.005 _minimize_neldermead  scipy/optimize/_optimize.py:708
|          |- 0.002 [self]  None
|          |- 0.002 function_wrapper  scipy/optimize/_optimize.py:564
|              |- 0.001 f  <string>:2
|                  |
|                  | |- 0.000 sum  <__array_function__ internals>:177
```

```

|         | | |  ^- 0.000 sum numpy/core/fromnumeric.py:2188
|         | | |      |- 0.000 _wrapreduction numpy/core/fromnumeric.py:69
|         | | |      |  |- 0.000 [self] None
|         | | |      |  |- 0.000 ufunc.reduce None
|         | | |      |  ^- 0.000 <dictcomp> numpy/core/fromnumeric.py:70
|         | | |      ^- 0.000 [self] None
|         | | ^- 0.000 [self] None
|         | ^- 0.000 copy < array function internals>:177

```



# optimize.minimize() output

```
1 f, grad, hess = mk_quad(0.7)
```

```
1 optimize.minimize(fun = f, x0 = (1.6, 1.1),  
2                   jac=grad, method="BFGS")
```

```
message: Optimization terminated successfully.  
success: True  
status: 0  
  fun: 1.2739256453438974e-11  
   x: [-5.318e-07 -8.843e-06]  
  nit: 6  
  jac: [-3.510e-07 -2.860e-06]  
hess_inv: [[ 1.515e+00 -3.438e-03]  
           [-3.438e-03  3.035e+00]]  
nfev: 7  
njev: 7
```

```
1 optimize.minimize(fun = f, x0 = (1.6, 1.1),  
2                   jac=grad, hess=hess,  
3                   method="Newton-CG")
```

```
message: Optimization terminated successfully.  
success: True  
status: 0  
  fun: 2.3418652989289317e-12  
   x: [ 0.000e+00  3.806e-06]  
  nit: 11  
  jac: [ 0.000e+00  4.102e-06]  
nfev: 12  
njev: 12  
nhev: 11
```

# optimize.minimize() output (cont.)

```
1 optimize.minimize(fun = f, x0 = (1.6, 1.1),  
2                   jac=grad, method="CG")
```

```
message: Optimization terminated successfully.  
success: True  
status: 0  
  fun: 1.4450021261144105e-32  
   x: [-1.943e-16 -1.110e-16]  
  nit: 2  
  jac: [-1.282e-16 -3.590e-17]  
 nfev: 5  
 njev: 5
```

```
1 optimize.minimize(fun = f, x0 = (1.6, 1.1),  
2                   jac=grad, method="Nelder-Mead")
```

```
message: Optimization terminated successfully.  
success: True  
status: 0  
  fun: 2.3077013477040082e-10  
   x: [ 1.088e-05  3.443e-05]  
  nit: 46  
 nfev: 89  
final_simplex: (array([[ 1.088e-05,  3.443e-05],  
                        [ 1.882e-05, -3.825e-05],  
                        [-3.966e-05, -3.147e-05]]), array([0.00000000e+00, 0.00000000e+00]))  
  
/opt/homebrew/lib/python3.10/site-packages/scipy/optimize/_minimize.py:100: UserWarning: Method Nelder-Mead does not use gradient information (jac=grad)
```

# Collect

```

1 def run_collect(name, x0, cost_func, *args, tol=
2     f, grad, hess = cost_func(*args)
3     methods = define_methods(x0, f, grad, hess, to
4
5     res = []
6     for method in methods:
7         if method in skip: continue
8
9         x = methods[method]()
10
11         d = {
12             "name":    name,
13             "method":  method,
14             "nit":     x["nit"],
15             "nfev":    x["nfev"],
16             "njev":    x.get("njev"),
17             "nhev":    x.get("nhev"),
18             "success": x["success"]
19             #"message": x["message"]
20         }
21         res.append( pd.DataFrame(d, index=[1]) )
22
23     return pd.concat(res)

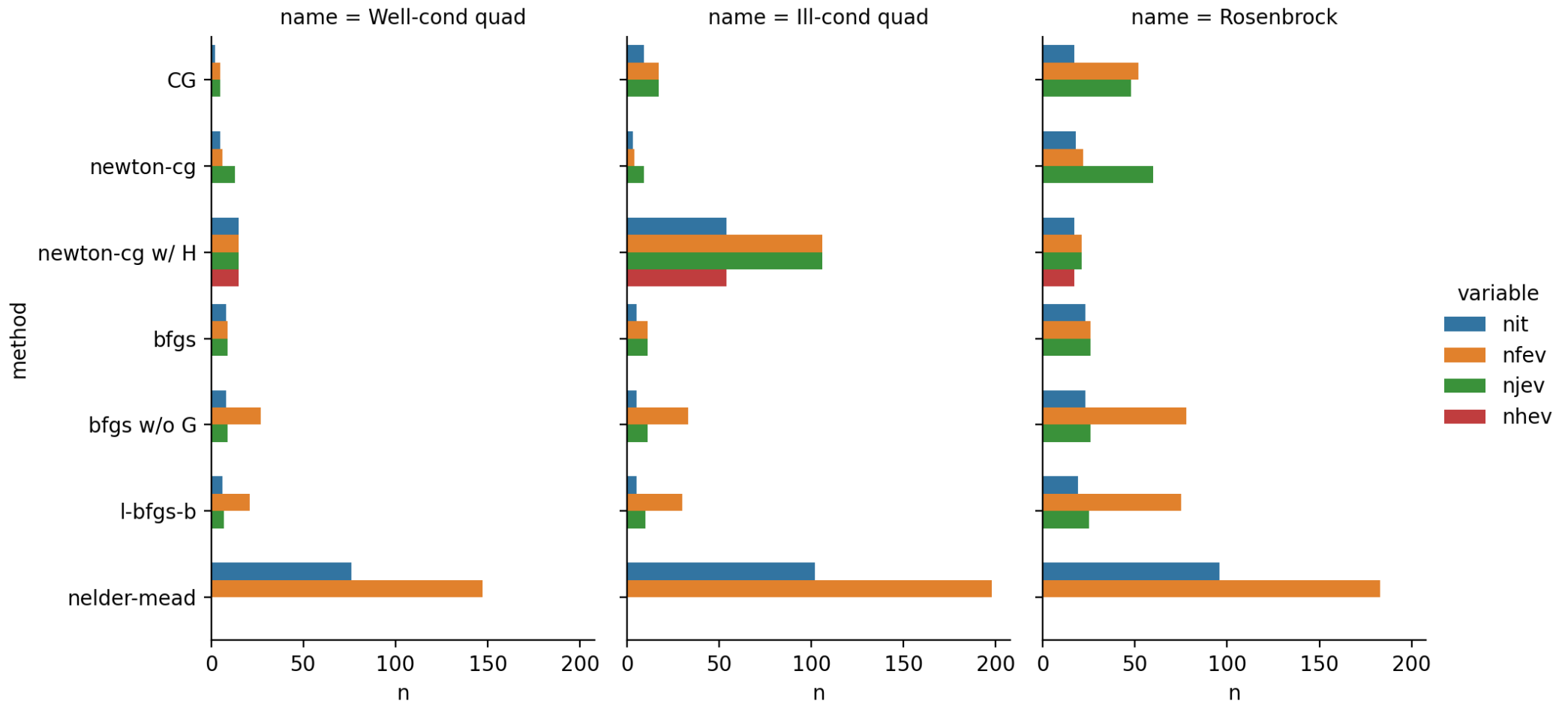
```

```

1 df = pd.concat([
2     run_collect(name, (1.6, 1.1), cost_func, arg,
3     for name, cost_func, arg in zip(
4         ("Well-cond quad", "Ill-cond quad", "Rosenbr
5         (mk_quad, mk_quad, mk_rosenbrock),
6         (0.7, 0.02, None)
7     )
8 ])
9
10 df

```

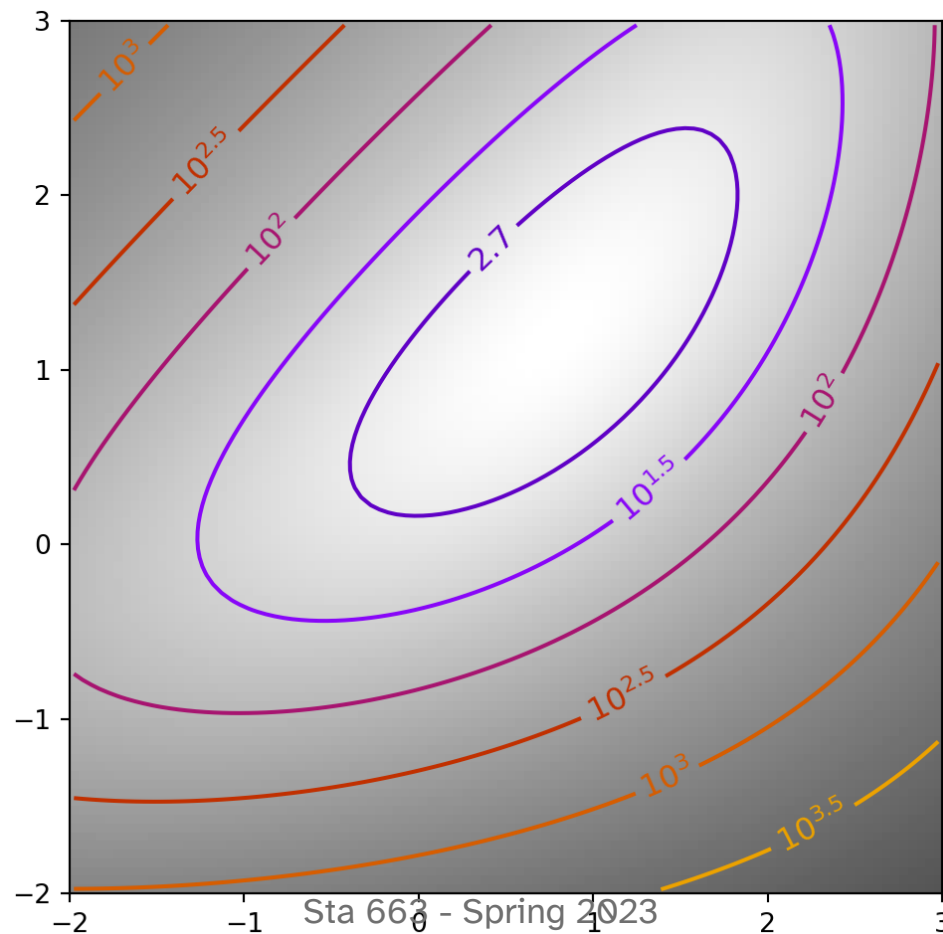
	name	method	nit	nfev	njev	r
1	Well-cond quad	CG	2	5	5	1
1	Well-cond quad	newton-cg	5	6	13	
1	Well-cond quad	newton-cg w/ H	15	15	15	
1	Well-cond quad	bfgs	8	9	9	1
1	Well-cond quad	bfgs w/o G	8	27	9	1
1	Well-cond quad	l-bfgs-b	6	21	7	1
1	Well-cond quad	nelder-mead	76	147	None	1
1	Ill-cond quad	CG	9	17	17	1
1	Ill-cond quad	newton-cg	3	4	9	
1	Ill-cond quad	newton-cg w/ H	54	106	106	
1	Ill-cond quad	bfgs	5	11	11	1
1	Ill-cond quad	bfgs w/o G	5	33	11	1
1	Ill-cond quad	l-bfgs-b	5	30	10	1
1	Ill-cond quad	nelder-mead	102	198	None	1
1	Rosenbrock	CG	17	52	48	1
1	Rosenbrock	newton-cg	18	22	60	
1	Rosenbrock	newton-cg w/ H	17	21	21	
1	Rosenbrock	bfgs	23	26	26	1
1	Rosenbrock	bfgs w/o G	23	78	26	1
1	Rosenbrock	l-bfgs-b	19	75	25	1



# Exercise 1

Try minimizing the following function using different optimization methods starting from  $\mathbf{x}_0 = [0, 0]$ , which method(s) appear to work best?

$$f(\mathbf{x}) = \exp(\mathbf{x}_1 - 1) + \exp(-\mathbf{x}_2 + 1) + (\mathbf{x}_1 - \mathbf{x}_2)^2$$



# MVN Example

# MVN density cost function

For an n-dimensional multivariate normal we define the  $n \times 1$  vectors  $\mathbf{x}$  and  $\boldsymbol{\mu}$  and the  $n \times n$  covariance matrix  $\boldsymbol{\Sigma}$ ,

$$f(\mathbf{x}) = \det(2\pi\boldsymbol{\Sigma})^{-1/2} \exp\left[-\frac{1}{2}(\mathbf{x} - \boldsymbol{\mu})^T \boldsymbol{\Sigma}^{-1} (\mathbf{x} - \boldsymbol{\mu})\right]$$

$$\nabla f(\mathbf{x}) = -f(\mathbf{x})\boldsymbol{\Sigma}^{-1}(\mathbf{x} - \boldsymbol{\mu})$$

$$\nabla^2 f(\mathbf{x}) = f(\mathbf{x}) (\boldsymbol{\Sigma}^{-1}(\mathbf{x} - \boldsymbol{\mu})(\mathbf{x} - \boldsymbol{\mu})^T \boldsymbol{\Sigma}^{-1} - \boldsymbol{\Sigma}^{-1})$$

Our goal will be to find the mode (maximum) of this density.

```
1 def mk_mvn(mu, Sigma):
2     Sigma_inv = np.linalg.inv(Sigma)
3     norm_const = 1 / (np.sqrt(np.linalg.det(2*np.pi*Sigma)))
4
5     # Returns the negative density (since we want
6     def f(x):
7         x_m = x - mu
8         return -(norm_const *
9                 np.exp(-0.5 * (x_m.T @ Sigma_inv @ x_m).i
10
11     def grad(x):
12         return (-f(x) * Sigma_inv @ (x - mu))
13
14     def hess(x):
15         n = len(x)
16         x_m = x - mu
17         return f(x) * ((Sigma_inv @ x_m).reshape((n,
18
19     return f, grad, hess
```



# Gradient checking

One of the most common issues when implementing an optimizer is to get the gradient calculation wrong which can produce problematic results. It is possible to numerically check the gradient function by comparing results between the gradient function and finite differences from the objective function via `optimize.check_grad()`.

```
1 # 2d
2 f, grad, hess = mk_mvn(np.zeros(2), np.eye(2,2))
3 optimize.check_grad(f, grad, np.zeros(2))
```

2.634178031930877e-09

```
1 optimize.check_grad(f, grad, np.ones(2))
```

5.213238144735062e-10

```
1 # 5d
2 f, grad, hess = mk_mvn(np.zeros(5), np.eye(5,5))
3 optimize.check_grad(f, grad, np.zeros(5))
```

2.6031257322754127e-10

```
1 optimize.check_grad(f, grad, np.ones(5))
```

1.725679820308689e-11

```
1 # 10d
2 f, grad, hess = mk_mvn(np.zeros(10), np.eye(10))
3 optimize.check_grad(f, grad, np.zeros(10))
```

2.8760747774580336e-12

```
1 optimize.check_grad(f, grad, np.ones(10))
```

2.850398669793798e-14

```
1 # 20d
2 f, grad, hess = mk_mvn(np.zeros(20), np.eye(20))
3 optimize.check_grad(f, grad, np.zeros(20))
```

4.965068306494546e-16

```
1 optimize.check_grad(f, grad, np.ones(20))
```

1.0342002372572841e-20

# Gradient checking (wrong gradient)

```
1 wrong_grad = lambda x: 2*grad(x)
```

```
1 # 2d
2 f, grad, hess = mk_mvn(np.zeros(2), np.eye(2,2))
3 optimize.check_grad(f, wrong_grad, [0,0])
```

2.634178031930877e-09

```
1 optimize.check_grad(f, wrong_grad, [1,1])
```

0.08280196633767578

```
1 # 5d
2 f, grad, hess = mk_mvn(np.zeros(5), np.eye(5))
3 optimize.check_grad(f, wrong_grad, np.zeros(5))
```

2.6031257322754127e-10

```
1 optimize.check_grad(f, wrong_grad, np.ones(5))
```

0.0018548087267515347

# Hessian checking

Note since the gradient of the gradient / jacobian is the hessian we can use this function to check our implementation of the hessian as well, just use `grad()` as `func` and `hess()` as `grad`.

```
1 # 2d
2 f, grad, hess = mk_mvn(np.zeros(2), np.eye(2,2))
3 optimize.check_grad(grad, hess, [0,0])
```

3.925231146709438e-17

```
1 optimize.check_grad(grad, hess, [1,1])
```

8.399162985270666e-10

```
1 # 5d
2 f, grad, hess = mk_mvn(np.zeros(5), np.eye(5))
3 optimize.check_grad(grad, hess, np.zeros(5))
```

3.878959614448864e-18

```
1 optimize.check_grad(grad, hess, np.ones(5))
```

3.8156075963144067e-11

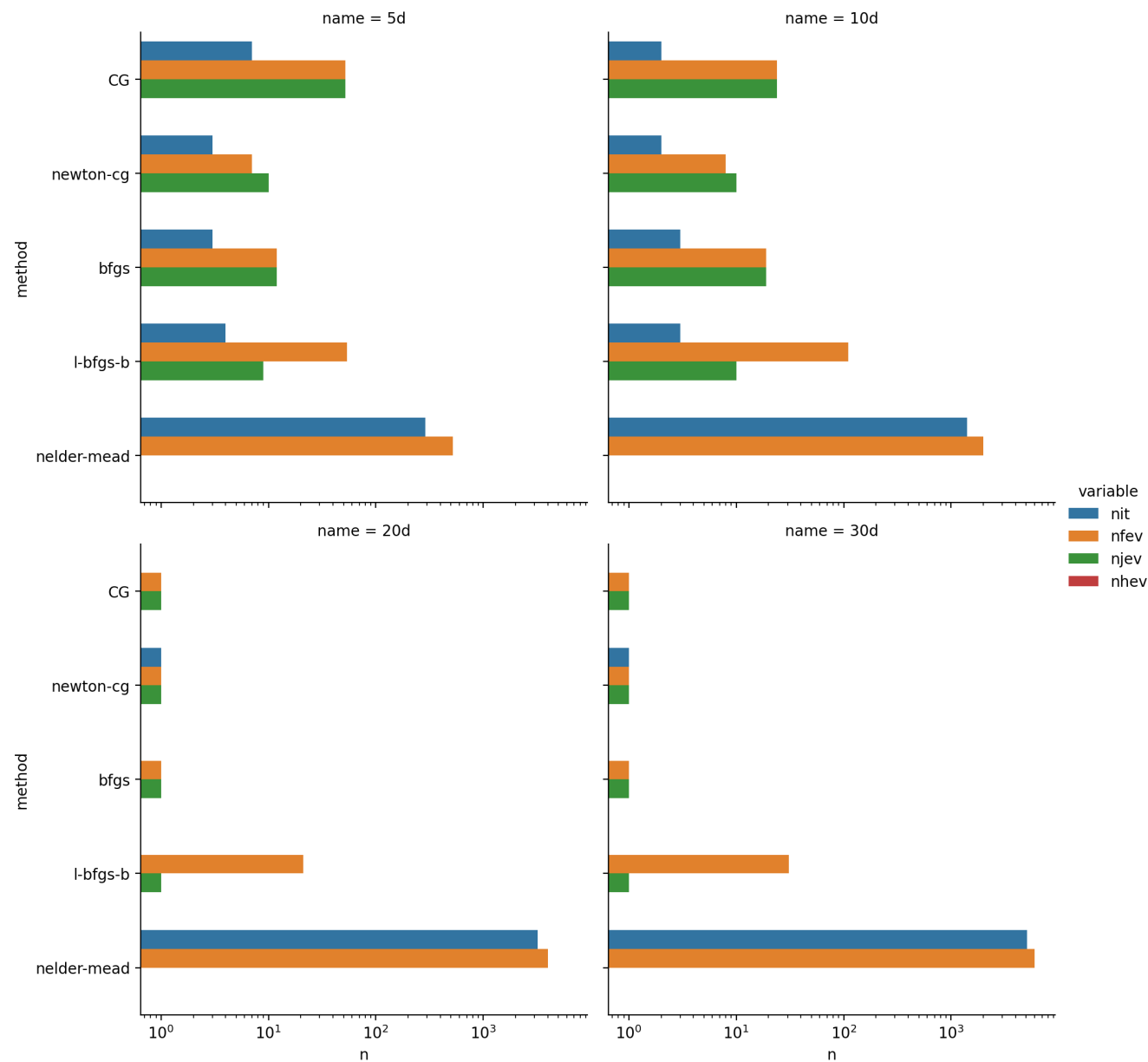
# Unit MVNs

```
1 df = pd.concat([
2     run_collect(
3         name, np.ones(n), mk_mvn,
4         np.zeros(n), np.eye(n),
5         tol=1e-10,
6         skip=['naive_newton', 'naive_cg', 'bfgs w/o
7     ])
8     for name, n in zip(
9         ("5d", "10d", "20d", "30d"),
10        (5, 10, 20, 30)
11    )
12 ])
```

```
1 df
```

	name	method	nit	nfev	njev	nhev	success
1	5d	CG	7	52	52	None	True
1	5d	newton-cg	3	7	10	0	True
1	5d	bfgs	3	12	12	None	True
1	5d	l-bfgs-b	4	54	9	None	True
1	5d	nelder-mead	290	523	None	None	True
1	10d	CG	2	24	24	None	True
1	10d	newton-cg	2	8	10	0	True
1	10d	bfgs	3	19	19	None	True
1	10d	l-bfgs-b	3	110	10	None	True
1	10d	nelder-mead	1403	2000	None	None	False
1	20d	CG	0	1	1	None	True
1	20d	newton-cg	1	1	1	0	True
1	20d	bfgs	0	1	1	None	True
1	20d	l-bfgs-b	0	21	1	None	True
1	20d	nelder-mead	3217	4000	None	None	False
1	30d	CG	0	1	1	None	True
1	30d	newton-cg	1	1	1	0	True
1	30d	bfgs	0	1	1	None	True
1	30d	l-bfgs-b	0	31	1	None	True
1	30d	nelder-mead	5097	6000	None	None	False

# Performance (Unit MVNs)

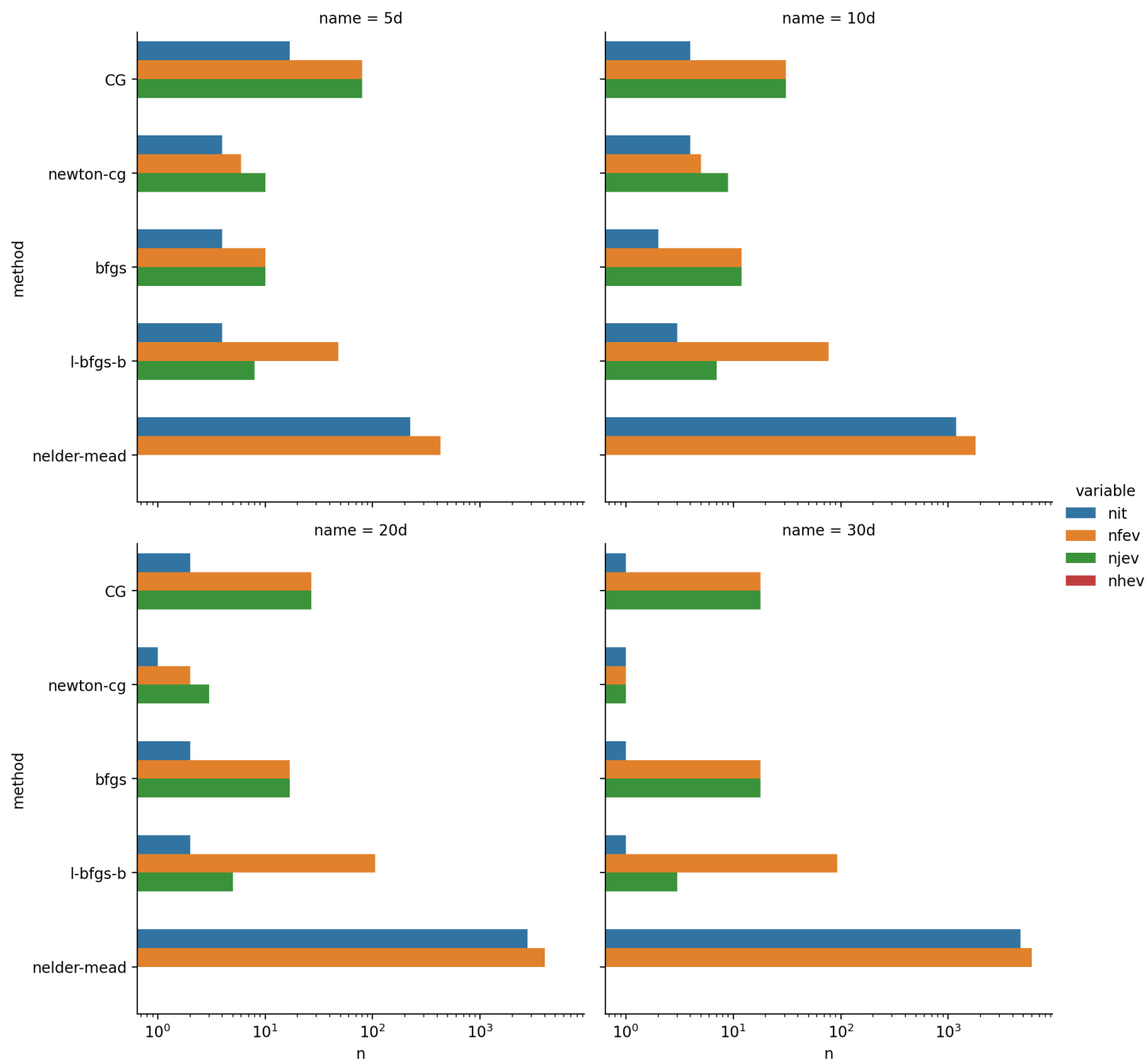


# Adding correlation

```
1 def build_Sigma(n, corr=0.5):
2     S = np.full((n,n), corr)
3     np.fill_diagonal(S, 1)
4     return S
5
6 df = pd.concat([
7     run_collect(
8         name, np.ones(n), mk_mvn,
9         np.zeros(n), build_Sigma(n),
10        tol=1e-10,
11        skip=['naive_newton', 'naive_cg', 'bfgs w/o
12    ])
13    for name, n in zip(
14        ("5d", "10d", "20d", "30d"),
15        (5, 10, 20, 30)
16    )
17 ])
```

```
1 df
```

	name	method	nit	nfev	njev	nhev	success
1	5d	CG	17	80	80	None	True
1	5d	newton-cg	4	6	10	0	True
1	5d	bfgs	4	10	10	None	True
1	5d	l-bfgs-b	4	48	8	None	True
1	5d	nelder-mead	224	427	None	None	True
1	10d	CG	4	31	31	None	True
1	10d	newton-cg	4	5	9	0	True
1	10d	bfgs	2	12	12	None	True
1	10d	l-bfgs-b	3	77	7	None	True
1	10d	nelder-mead	1184	1802	None	None	True
1	20d	CG	2	27	27	None	True
1	20d	newton-cg	1	2	3	0	True
1	20d	bfgs	2	17	17	None	True
1	20d	l-bfgs-b	2	105	5	None	True
1	20d	nelder-mead	2745	4000	None	None	False
1	30d	CG	1	18	18	None	True
1	30d	newton-cg	1	1	1	0	True
1	30d	bfgs	1	18	18	None	True
1	30d	l-bfgs-b	1	93	3	None	True
1	30d	nelder-mead	4687	6000	None	None	False



# What's going on? (good)

```
1 n = 5
2 f, grad, hess = mk_mvn(np.zeros(n), build_Sigma(n))
```

```
1 optimize.minimize(f, np.ones(n), jac=grad,
2                   method="CG", tol=1e-9)
```

```
message: Optimization terminated successfully.
success: True
status: 0
  fun: -0.023337250777292103
   x: [ 1.086e-07  1.061e-07  1.080e-07  1.080e-07]
  nit: 14
  jac: [ 8.802e-10  7.646e-10  8.540e-10  8.532e-10]
nfev: 67
njev: 67
```

```
1 optimize.minimize(f, np.ones(n), jac=grad,
2                   method="CG", tol=1e-10)
```

```
message: Optimization terminated successfully.
success: True
status: 0
  fun: -0.023337250777292328
   x: [-2.232e-09 -3.779e-10 -1.811e-09 -1.797e-09]
  nit: 17
  jac: [-4.415e-11  4.237e-11 -2.453e-11 -2.387e-11]
nfev: 80
njev: 80
```



```
1 optimize.minimize(f, np.ones(n), jac=grad,  
2                   method="CG", tol=1e-11)
```

```
message: Desired error not necessarily achieved due  
success: False  
status: 2  
  fun: -0.02333725077729232  
   x: [ 2.205e-08  3.734e-09  1.790e-08  1.776e-08]  
  nit: 16  
  jac: [ 4.362e-10 -4.186e-10  2.424e-10  2.359e-10]  
 nfev: 93  
 njev: 81
```

# What's going on? (okay)

```
1 n = 20
2 f, grad, hess = mk_mvn(np.zeros(n), build_Sigma(n))
```

```
1 optimize.minimize(f, np.ones(n), jac=grad,
2                   method="CG", tol=1e-9)
```

```
message: Optimization terminated successfully.
success: True
status: 0
  fun: -2.330191334018497e-06
    x: [-3.221e-04 -3.221e-04 ... -3.221e-04 -3.221e-04]
  nit: 2
  jac: [-7.148e-11 -7.148e-11 ... -7.148e-11 -7.148e-11]
nfev: 27
njev: 27
```

```
1 optimize.minimize(f, np.ones(n), jac=grad,
2                   method="CG", tol=1e-10)
```

```
message: Optimization terminated successfully.
success: True
status: 0
  fun: -2.330191334018497e-06
    x: [-3.221e-04 -3.221e-04 ... -3.221e-04 -3.221e-04]
  nit: 2
  jac: [-7.148e-11 -7.148e-11 ... -7.148e-11 -7.148e-11]
nfev: 27
njev: 27
```

```
1 optimize.minimize(f, np.ones(n), jac=grad,  
2                   method="CG", tol=1e-11)
```

message: Optimization terminated successfully.

success: True

status: 0

fun: -2.3301915597315495e-06

x: [-4.506e-05 -4.506e-05 ... -4.506e-05 -4.506e-05]

nit: 2884

jac: [-9.999e-12 -9.999e-12 ... -9.999e-12 -9.999e-12]

nfev: 66313

njev: 66313

# What's going on? (bad)

```
1 n = 30
2 f, grad, hess = mk_mvn(np.zeros(n), build_Sigma(n))
```

```
1 optimize.minimize(f, np.ones(n), jac=grad,
2                   method="CG", tol=1e-9)
```

```
message: Optimization terminated successfully.
success: True
status: 0
  fun: -2.3811463025973114e-09
   x: [ 1.000e+00  1.000e+00 ...  1.000e+00  1.000e+00]
  nit: 0
  jac: [ 1.536e-10  1.536e-10 ...  1.536e-10  1.536e-10]
nfev: 1
njev: 1
```

```
1 optimize.minimize(f, np.ones(n), jac=grad,
2                   method="CG", tol=1e-10)
```

```
message: Optimization terminated successfully.
success: True
status: 0
  fun: -6.180056227752818e-09
   x: [ 1.203e-01  1.203e-01 ...  1.203e-01  1.203e-01]
  nit: 1
  jac: [ 4.795e-11  4.795e-11 ...  4.795e-11  4.795e-11]
nfev: 18
njev: 18
```

```
1 optimize.minimize(f, np.ones(n), jac=grad,  
2                   method="CG", tol=1e-11)
```

message: Optimization terminated successfully.

success: True

status: 0

fun: -6.26701117075865e-09

x: [-5.021e-03 -5.021e-03 ... -5.021e-03 -5.021e-03]

nit: 2

jac: [-2.030e-12 -2.030e-12 ... -2.030e-12 -2.030e-12]

nfev: 35

njev: 35

# Options (bfgs)

```
1 optimize.show_options(solver="minimize", method="bfgs")
```

Minimization of scalar function of one or more variables using the BFGS algorithm.

## Options

-----

`disp` : bool

Set to True to print convergence messages.

`maxiter` : int

Maximum number of iterations to perform.

`gtol` : float

Terminate successfully if gradient norm is less than ``gtol``.

`norm` : float

Order of norm (Inf is max, -Inf is min).

`eps` : float or ndarray

If ``jac`` is None` the absolute step size used for numerical approximation of the jacobian via forward differences.

`return_all` : bool, optional

Set to True to return a list of the best solution at each of the iterations.

`finite_diff_rel_step` : None or array\_like, optional

If ``jac`` in ['2-point', '3-point', 'cs']` the relative step size to use for numerical approximation of the jacobian. The absolute step size is computed as ``h` = rel_step * sign(x) * max(1, abs(x))``,

# Options (Nelder-Mead)

```
1 optimize.show_options(solver="minimize", method="Nelder-Mead")
```

Minimization of scalar function of one or more variables using the Nelder-Mead algorithm.

Options

-----

disp : bool

Set to True to print convergence messages.

maxiter, maxfev : int

Maximum allowed number of iterations and function evaluations.  
Will default to ``N\*200``, where ``N`` is the number of variables, if neither ``maxiter`` or ``maxfev`` is set. If both ``maxiter`` and ``maxfev`` are set, minimization will stop at the first reached.

return\_all : bool, optional

Set to True to return a list of the best solution at each of the iterations.

initial\_simplex : array\_like of shape (N + 1, N)

Initial simplex. If given, overrides ``x0``.

``initial\_simplex[j,:]`` should contain the coordinates of the jth vertex of the ``N+1`` vertices in the simplex, where ``N`` is the dimension.

xatol : float, optional

Absolute error in xopt between iterations that is acceptable for

# SciPy implementation

The following code comes from SciPy's `minimize()` implementation:

```
1  if tol is not None:
2      options = dict(options)
3      if meth == 'nelder-mead':
4          options.setdefault('xatol', tol)
5          options.setdefault('fatol', tol)
6      if meth in ('newton-cg', 'powell', 'tnc'):
7          options.setdefault('xtol', tol)
8      if meth in ('powell', 'l-bfgs-b', 'tnc', 'slsqp'):
9          options.setdefault('ftol', tol)
10     if meth in ('bfgs', 'cg', 'l-bfgs-b', 'tnc', 'dogleg',
11                'trust-ncg', 'trust-exact', 'trust-krylov'):
12         options.setdefault('gtol', tol)
13     if meth in ('cobyla', '_custom'):
14         options.setdefault('tol', tol)
15     if meth == 'trust-constr':
16         options.setdefault('xtol', tol)
17         options.setdefault('gtol', tol)
18         options.setdefault('barrier_tol', tol)
```



# Some general advice

- Having access to the gradient is almost always helpful / necessary
- Having access to the hessian can be helpful, but usually does not significantly improve things
- The curse of dimensionality is real
  - Be careful with `tol` - it means different things for different methods
- In general, **BFGS** or **L-BFGS** should be a first choice for most problems (either well- or ill-conditioned)
  - **CG** can perform better for well-conditioned problems with cheap function evaluations

# Maximum Likelihood example

# Normal MLE

```
1 from scipy.stats import norm
2
3 n = norm(-3.2, 1.25)
4 x = n.rvs(size=100, random_state=1234)
5 x.round(2)
```

```
array([-2.61, -4.69, -1.41, -3.59, -4.1 , -2.09, -2.1 ,
        -6.   , -1.76, -1.96, -2.01, -5.73, -3.62, -3.2 ,
        -1.55, -5.13, -3.45, -4.02, -2.96, -2.51, -1.5 ,
        -5.47, -3.43, -1.88, -3.7 , -2.78, -1.89, -1.8 ,
        -3.04, -3.6 , -2.15, -0.21, -3.1 , -3.91, -3.3 ,
        -4.32, -3.37, -3.18, -2.26, -2.93, -2.15, -5.0 ,
        -3.89, -3.38, -2.76, -3.24, -2.49, -1.27, -4.4 ,
        -3.46, -1.91, -6.2 , -0.66, -4.63, -2.94, -2.3 ,
        -2.32, -2.55, -4.36, -0.69, -2.92, -4.64, -2.4 ,
        -7.65, -1.55, -3.01, -2.99, -3.74, -2.24, -1.9 ,
        -3.1 , -3.7 , -4.48, -3.93, -2.18, -3.3 , -3.0 ,
        -3.84])
```

```
1 {'μ': x.mean(), 'σ': x.std()}
```

```
{'μ': -3.156109646093205, 'σ': 1.2446060629192535}
```

```
1 mle_norm = lambda θ: -np.sum(
2     norm.logpdf(x, loc=θ[0], scale=θ[1])
3 )
4
5 mle_norm([0,1])
```

```
667.3974708213642
```

```
1 mle_norm([-3, 1])
```

```
170.56457699340282
```

```
1 mle_norm([-3.2, 1.25])
```

```
163.83926813257395
```

# Minimizing

```
1 optimize.minimize(mle_norm, x0=[0,1], method="bfgs")
```

```
message: Desired error not necessarily achieved due to precision loss.
```

```
success: False
```

```
status: 2
```

```
fun: nan
```

```
x: [-1.436e+04 -3.533e+03]
```

```
nit: 2
```

```
jac: [      nan      nan]
```

```
hess_inv: [[ 9.443e-01  2.340e-01]
            [ 2.340e-01  5.905e-02]]
```

```
nfev: 339
```

```
njev: 113
```

# Adding constraints

```
1 def mle_norm2(theta):  
2     if theta[1] <= 0:  
3         return np.Inf  
4     else:  
5         return -np.sum(norm.logpdf(x, loc=theta[0], scale=theta[1]))
```

```
1 optimize.minimize(mle_norm2, x0=[0,1], method="bfgs")
```

message: Optimization terminated successfully.

success: True

status: 0

fun: 163.77575977255518

x: [-3.156e+00 1.245e+00]

nit: 9

jac: [-1.907e-06 0.000e+00]

hess\_inv: [[ 1.475e-02 -1.069e-04]

[-1.069e-04 7.734e-03]]

nfev: 47

njev: 15

/opt/homebrew/lib/python3.10/site-packages/scipy/optimize/\_minimize.py:100: RuntimeWarning: divide by zero encountered in divide

df = fun(x) - f0

# Specifying Bounds

It is also possible to specify bounds via `bounds` but this is only available for certain optimization methods.

```
1 optimize.minimize(  
2     mle_norm, x0=[0,1], method="l-bfgs-b",  
3     bounds = [(-1e16, 1e16), (1e-16, 1e16)]  
4 )
```

```
message: CONVERGENCE: REL_REDUCTION_OF_F_<=_FACTR*EPSMCH  
success: True  
status: 0  
  fun: 163.7757597728758  
   x: [-3.156e+00  1.245e+00]  
  nit: 10  
  jac: [ 2.075e-04  0.000e+00]  
 nfev: 69  
 njev: 23  
hess_inv: <2x2 LbfgsInvHessProduct with dtype=float64>
```

# Exercise 2

Using `optimize.minimize()` recover the shape and scale parameters for these data using MLE.

```
1 from scipy.stats import gamma
2
3 g = gamma(a=2.0, scale=2.0)
4 x = g.rvs(size=100, random_state=1234)
5 x.round(2)
```

```
array([ 4.7 ,  1.11,  1.8 ,  6.19,  3.37,  0.25,  6.45,  0.36,  4.49,
        4.14,  2.84,  1.91,  8.03,  2.26,  2.88,  6.88,  6.84,  6.83,
        6.1 ,  3.03,  3.67,  2.57,  3.53,  2.07,  4.01,  1.51,  5.69,
        3.92,  6.01,  0.82,  2.11,  2.97,  5.02,  9.13,  4.19,  2.82,
       11.81,  1.17,  1.69,  4.67,  1.47, 11.67,  5.25,  3.44,  8.04,
        3.74,  5.73,  6.58,  3.54,  2.4 ,  1.32,  2.04,  2.52,  4.89,
        4.14,  5.02,  4.75,  8.24,  7.6 ,  1.   ,  6.14,  0.58,  2.83,
        2.88,  5.42,  0.5 ,  3.46,  4.46,  1.86,  4.59,  2.24,  2.62,
        3.99,  3.74,  5.27,  1.42,  0.56,  7.54,  5.5 ,  1.58,  5.49,
        6.57,  4.79,  5.84,  8.21,  1.66,  1.53,  4.27,  2.57,  1.48,
        5.23,  3.84,  3.15,  2.1 ,  3.71,  2.79,  0.86,  8.52,  4.36,
        3.3 ])
```

