STAT 4830: Numerical optimization for data science and ML

Lecture 3: Linear Regression - Gradient Descent

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The Memory Wall

Consider genomic prediction: 1000 patients, 100,000 genetic markers

```
n_samples = 1000
n_markers = 100_000
memory_needed = (n_markers * n_markers * 8) / (1024**3) # in GB
print(f"Memory needed for X^TX: {memory_needed:.1f} GB") # 80.0 GB
```

Just forming $X^ op X$ would exceed most workstations' memory!

Even Worse: Medical Imaging

MRI reconstruction with 256^3 voxels:

- Matrix size: $256^3 imes 256^3$
- ullet Memory for $X^ op X$: 2.2 petabytes
- That's 0.2% of world's total data center storage in 2023!

These aren't edge cases - they're routine analysis tasks.

Why Direct Methods Fail

Direct methods solve normal equations $X^ op Xw = X^ op y$:

```
# Direct method (fails for large p)
XtX = X.T @ X  # Form p × p matrix
Xty = X.T @ y  # Form p × 1 vector
w = solve(XtX, Xty)  # Solve p × p system
```

Costs:

- 1. Forming $X^ op X$: $O(np^2)$ operations, $O(p^2)$ memory
- 2. Forming $X^ op y$: O(np) operations, O(p) memory
- 3. Solving system: $O(p^3)$ operations

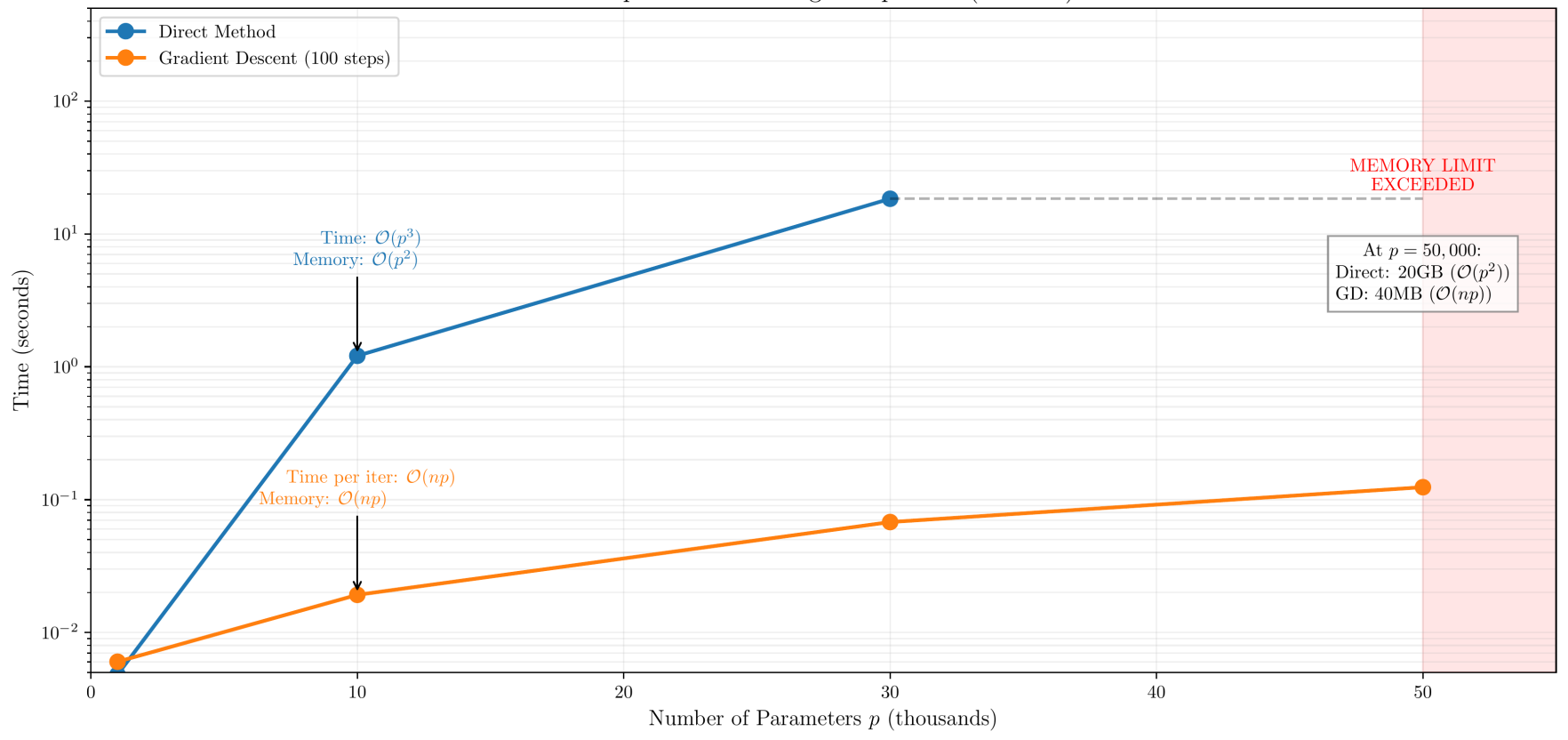
Experimental Results: Memory Wall

Results on MacBook M1 Pro (64GB RAM):

```
Size (p)
           Memory for X^TX
                           Time
                                      Status
1,000
           8MB
                            0.005s
                                     Fast, fits in fast memory
5,000
                           0.182s Fits in RAM
           200MB
       3.2GB
20,000
                           5.209s RAM stressed
50,000
           20GB
                           FAILS
                                    Out of memory
```

Memory becomes bottleneck before computation time!

Experimental Results: Scaling Behavior



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The pattern is clear: memory becomes the bottleneck long before computation time:

A Memory-Efficient Alternative

One memory-efficient alternative is gradient descent:

```
# This forms a huge p × p matrix (bad)
XtX = X.T @ X  # Need O(p²) memory
result = XtX @ w  # Matrix-vector product

# Gradient descent uses operations like these:
Xw = X @ w  # Need O(p) memory
```

Both compute $(X^ op X)w$, but gradient descent:

- ullet Never forms the p imes p matrix
- Uses O(np) operations (same as first approach)
- ullet Only needs O(p) extra memory for vectors

The Algorithm

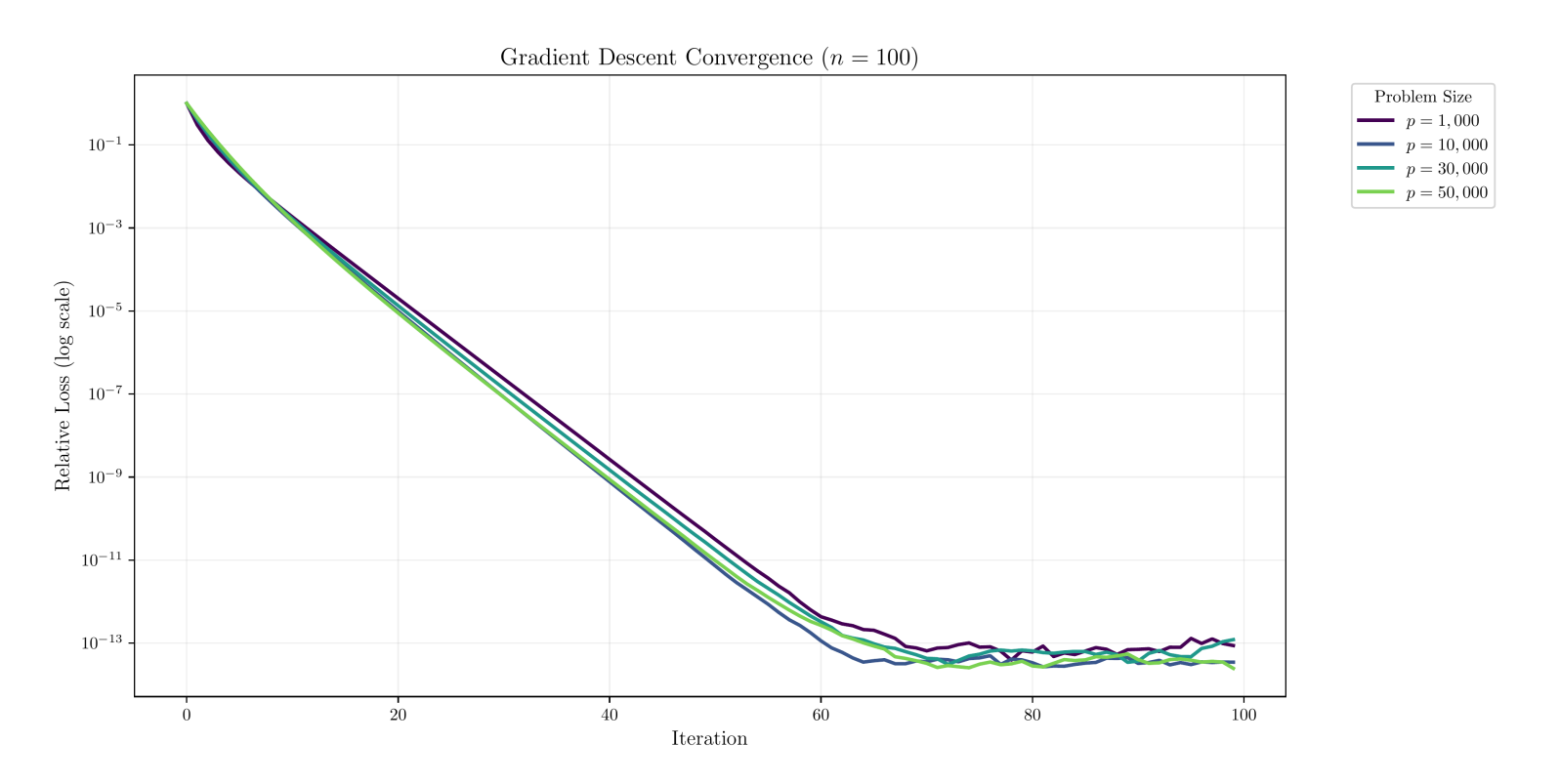
```
# Gradient descent with matrix-vector products
w = torch.zeros(p)  # Initial guess
for k in range(max_iters):
    Xw = X @ w  # Forward pass: O(np)
    grad = X.T @ (Xw - y)  # Backward pass: O(np)
    w -= step_size * grad # Update: O(p)
```

The memory efficiency comes from iteratively updating our solution:

- 1. Start with an initial guess (even all zeros)
- 2. Compute the gradient using matrix-vector products
- 3. Take a small step in that direction
- 4. Repeat until convergence

Convergence Behavior: The Pattern

Our experiments with random matrices reveal a fascinating pattern:



Convergence Behavior: Key Insights

Linear Convergence

Error decreases exponentially, appearing as a straight line on log scale. This predictable rate of improvement lets us estimate progress.

Precision vs Time

Each doubling of iterations improves precision by $\sim\!10^4$. This consistent behavior lets us plan computational resources.

Practical Impact

- 20 iterations: $\sim 10^{-5}$ relative error
- 40 iterations: $\sim 10^{-9}$ relative error
- 60 iterations: $\sim 10^{-13}$ relative error

The Least Squares Landscape

Our objective measures squared prediction error:

$$f(w) = rac{1}{2} \|Xw - y\|_2^2 = rac{1}{2} (Xw - y)^ op (Xw - y)$$

Expanding reveals the quadratic structure:

$$f(w) = rac{1}{2}(w^ op X^ op Xw - 2y^ op Xw + y^ op y)$$

Each term has meaning:

- $ullet w^ op X^ op Xw = \|Xw\|^2$: size of predictions
- ullet $2y^ op Xw$: alignment with truth
- $y^{\top}y$: scale of target values

Computing the Gradient

The gradient has a beautiful form:

$$egin{aligned} rac{\partial f}{\partial w_j} &= \sum_{i=1}^n (x_i^ op w - y_i) x_{ij} \
abla f(w) &= X^ op (Xw - y) = X^ op Xw - X^ op y \end{aligned}$$

This tells us:

- ullet Xw-y is prediction error in output space
- ullet $X^ op$ projects error back to parameter space
- Direction tells us how to adjust each parameter

Finding the Direction of Steepest Descent

For our quadratic function, we can compute the exact change:

$$egin{aligned} f(w+\epsilon v) &= rac{1}{2}\|X(w+\epsilon v)-y\|_2^2 \ &= f(w) + \epsilon (Xw-y)^ op Xv + rac{\epsilon^2}{2}v^ op X^ op Xv \ &= f(w) + \epsilon
abla f(w)^ op v + rac{\epsilon^2}{2}v^ op X^ op Xv \end{aligned}$$

For small ϵ , the ϵ dominates ϵ^2 .

Linear Approximation

IDEA: At any point w, we can approximate f using its gradient:

$$f(w + \epsilon v) pprox f(w) + \epsilon
abla f(w)^ op v$$

This so-called **first-order approximation**:

- Determines initial rate of descent
- Guides stepsize selection
- Explains convergence behavior

The Optimization Problem

At any point w, we want the direction v that decreases the first order approximation of f most rapidly:

minimize
$$abla f(w)^ op v$$
 subject to $\|v\|=1$

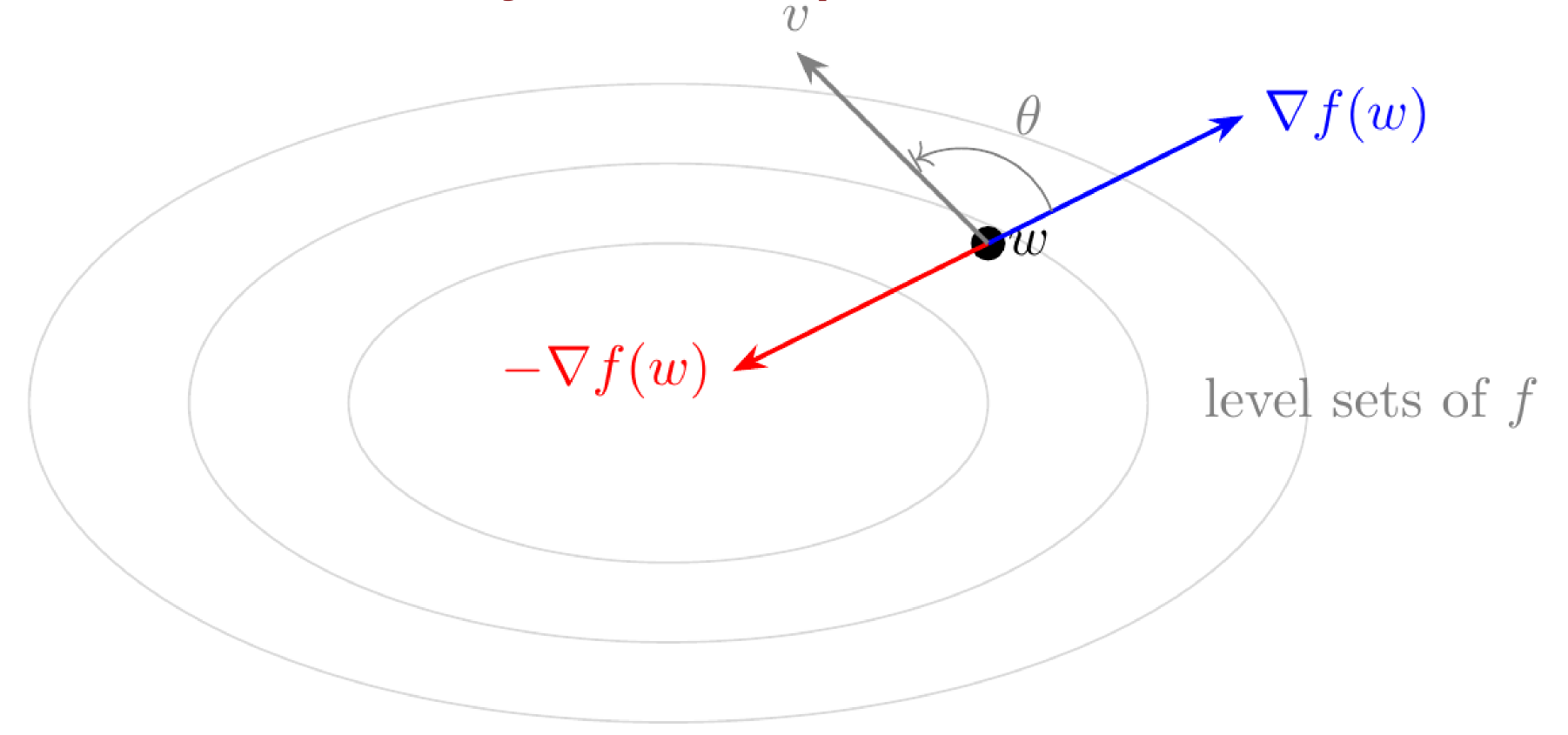
The solution is:

$$v_{\star} = -rac{
abla f(w)}{\|
abla f(w)\|}$$

Indeed, by Cauchy-Schwarz inequality:

$$|
abla f(w)^ op v| \leq \|
abla f(w)\| \|v\| = \|
abla f(w)\|$$

The Geometry of Steepest Descent



What if the Gradient is zero?

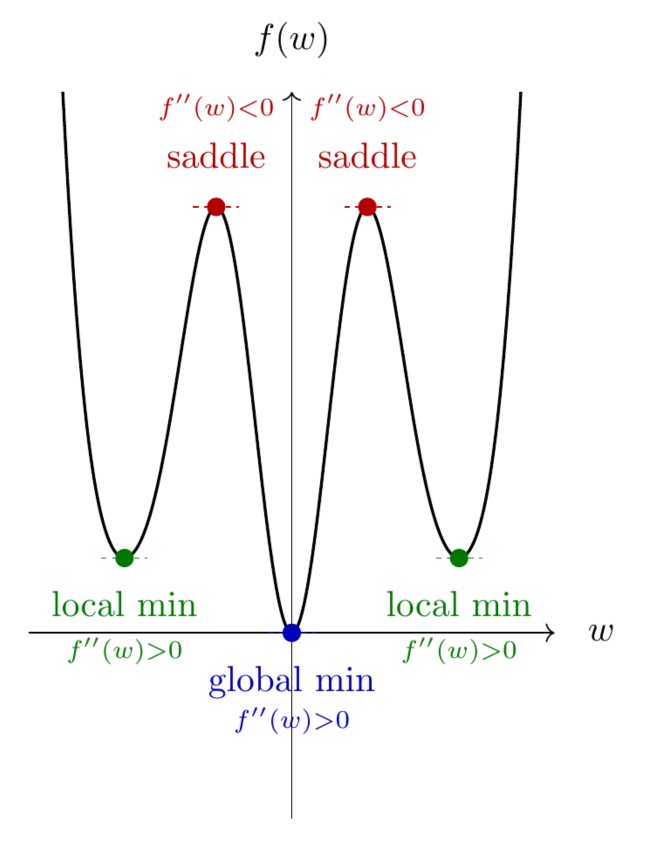
When abla f(w) = 0, we've found a critical point:

- Local Minimum: All directions curve upward
- Local Maximum: All directions curve downward
- Saddle Point: Some up, some down

For least squares: All critical points are global minima!

• This is due to **convexity** -- a property we'll study later.

What if the Gradient is zero?



The Algorithm: Overview

At each step:

- 1. Start at our current point w_k
- 2. Compute the gradient $g_k = X^ op X w_k X^ op y$
- 3. Move in the negative gradient direction: $w_{k+1} = w_k lpha_k g_k$
- 4. Repeat until the gradient becomes small

Three key factors determine success:

- Stepsize selection
- Problem conditioning
- Initial guess quality

The Algorithm: Implementation

```
# Gradient descent with matrix-vector products
w = torch.zeros(p)  # Initial guess
for k in range(max_iters):
    Xw = X @ w  # Forward pass: O(np)
    grad = X.T @ (Xw - y)  # Backward pass: O(np)
    w -= step_size * grad # Update: O(p)
```

For least squares, starting at zero is reasonable:

- Gives zero predictions a natural baseline
- Will eventually find minimum (thanks to convexity)
- Good initial guess reduces iterations needed

Stepsize Selection: The Theory

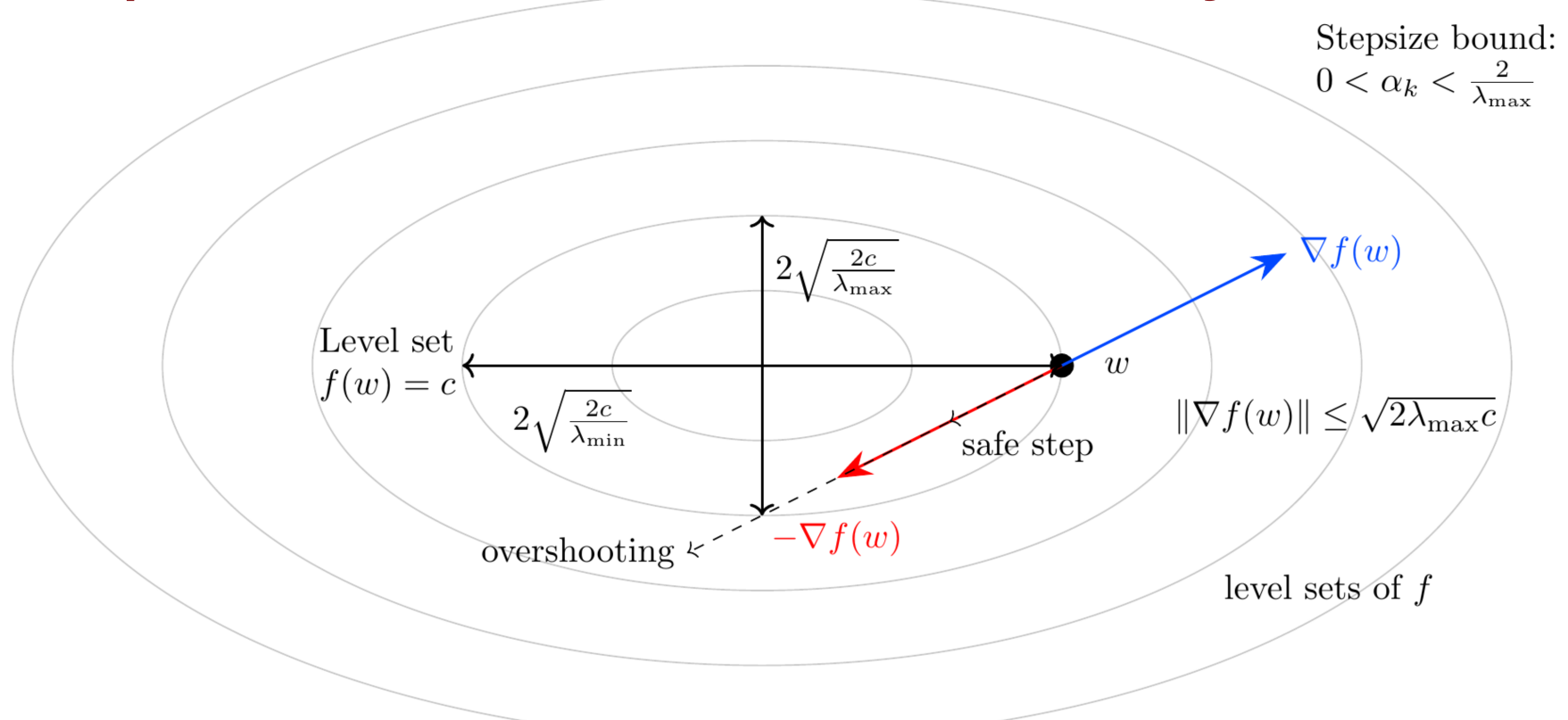
Convergence is guaranteed when:

$$0 < lpha_k < rac{2}{\lambda_{ ext{max}}(X^ op X)}$$

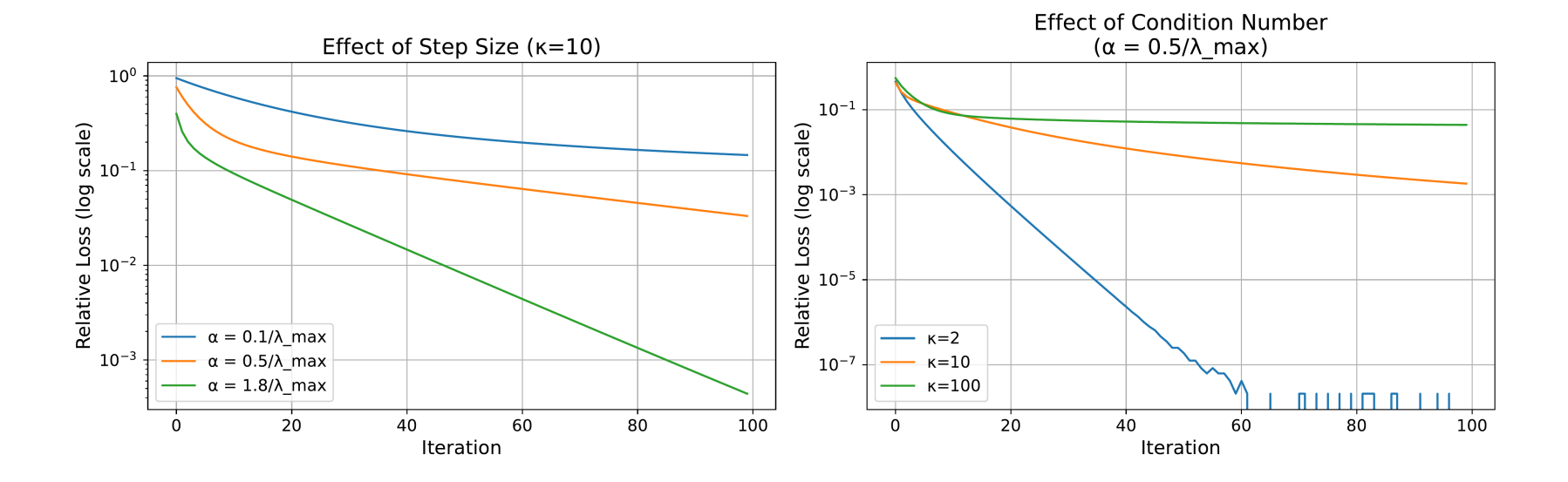
Why this bound?

- Level sets become very narrow in some directions
- ullet Width determined by eigenvalues of $X^ op X$
- Too large a step overshoots the minimum

Stepsize Selection: The Geometry



Convergence Speed vs Condition Number



Left: Effect of stepsize ($\kappa=10$)

Right: Effect of condition number (fixed stepsize)

Effect of Condition Number: Analysis

The path to the minimum depends on problem conditioning:

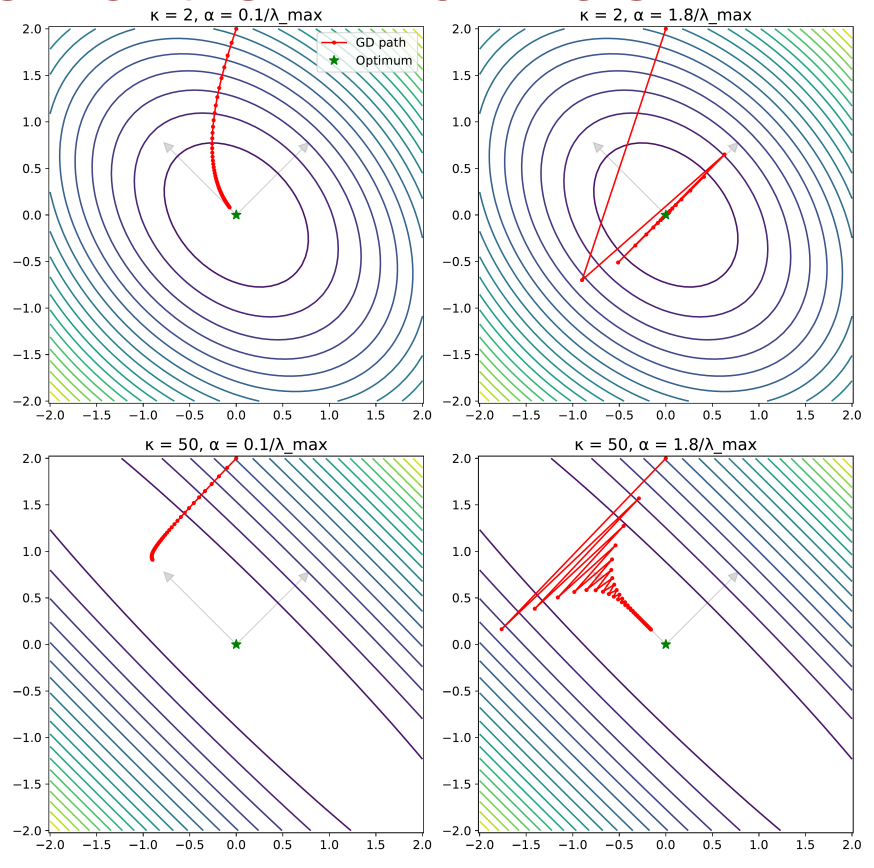
Well-Conditioned ($\kappa=2$)

- Direct path to minimum
- Fast, steady progress
- Efficient use of computation

Poorly-Conditioned ($\kappa=50$)

- Zigzag path to minimum
- Slow overall progress
- Many wasted steps

Effect of Condition Number $\kappa = 2, \alpha = 0.1/\lambda_{\text{max}}$ $\kappa = 2, \alpha = 1.8/\lambda_{\text{max}}$



Limitations and Next Steps

Gradient descent also has limitations:

- ullet For large n: Computing full gradient expensive
- ullet For large p: Memory still scales with problem size
- Poor conditioning: Slow convergence

Solutions we'll cover later:

- 1. Stochastic methods for large n
- 2. Coordinate descent for large p
- 3. Momentum and adaptive methods for conditioning

Summary

- 1. Memory Wall: Direct methods fail for large problems
- 2. Gradient Descent: Memory-efficient iterative solution
- 3. Convergence: Linear rate with predictable behavior
- 4. Geometry: Follows steepest descent direction
- 5. Implementation: Simple, scalable algorithm
- 6. Limitations: Sets up need for advanced methods

Next lecture: problems beyond least squares.