# Large-Scale Derivative-Free Optimization using Subspace Methods

Joint work with Coralia Cartis (Oxford)

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### Outline

- 1. Scalability of model-based DFO
- 2. Subspace DFO methods: algorithm & theory
- 3. Specialization to least-squares: numerical results

### Model-Based DFO — Basic Ideas

### Model-Based DFO

$$\min_{\mathbf{x}\in\mathbb{R}^n}f(\mathbf{x})$$

• Classically (e.g. Newton's method),

$$f(\mathbf{x}_k + \mathbf{s}) \approx m_k(\mathbf{s}) = f(\mathbf{x}_k) + \nabla f(\mathbf{x}_k)^T \mathbf{s} + \frac{1}{2} \mathbf{s}^T \nabla^2 f(\mathbf{x}_k) \mathbf{s}$$

Instead, build interpolation model

$$f(\mathbf{x}_k + \mathbf{s}) \approx m_k(\mathbf{s}) = f(\mathbf{x}_k) + \mathbf{g}_k^T \mathbf{s} + \frac{1}{2} \mathbf{s}^T \mathbf{H}_k \mathbf{s}$$

- ullet Geometry of points good  $\Longrightarrow$  interpolation model Taylor-accurate  $\Longrightarrow$  convergence
- Global convergence via trust-region method

[Powell, 2003; Conn, Scheinberg & Vicente, 2009]

### Model-Based DFO — Theory

Model-based methods have similar convergence results to derivative-based methods.

**Worst-case complexity:** how many iterations before  $\epsilon$  accuracy guaranteed?

Accuracy order	Model-based DFO	Taylor models
1st: $\ \nabla f(\mathbf{x}_k)\ _2 \leq \epsilon$	$\mathcal{O}(n^2\epsilon^{-2})$	$\mathcal{O}(\epsilon^{-2})$
2nd: 1st & $\lambda_{\min}(\nabla^2 f(\boldsymbol{x}_k)) \geq -\epsilon$	$\mathcal{O}(n^9\epsilon^{-3})$	$\mathcal{O}(\epsilon^{-3})$

[Cartis, Gould & Toint, 2010; Garmanjani, Júdice & Vicente, 2016]

- ullet Same  $\epsilon$  dependency as derivative-based, but scales badly with problem dimension n
- Substantial linear algebra work for interpolation and geometry management:
  - $\mathcal{O}(n^3)$  flops per iteration for linear models,  $\mathcal{O}(n^6)$  for quadratic models.

### Challenge

How can DFO methods be made scalable?

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#### Scalable DFO

### Challenge

How can DFO methods be made scalable?

- Exploit known problem structure [Porcelli & Toint, 2020; Bandeira et al., 2012]
- Randomized finite differencing ('gradient sampling')
   [Nesterov & Spokoiny, 2017]
- Randomized direct search: sample a subset of search directions improves complexity from  $\mathcal{O}(n^2\epsilon^{-2})$  to  $\mathcal{O}(n\epsilon^{-2})$  [Gratton et al., 2015; Bergou et al., 2020]

### Applications for scalable DFO methods include:

- Machine learning [Salimans et al., 2017; Ughi et al., 2020]
- Image analysis [Ehrhardt & R., 2021]
- Proxy for global optimization methods [Cartis, R. & Sheridan-Methven, 2021]

# **Subspace DFO**

We use a subspace method: only search in low-dimensional subspaces of  $\mathbb{R}^n$ 

• Related to coordinate descent methods

- [Wright, 2015; Patrascu & Necoara, 2015]
- Some implementations exist, but no theory [Gross & Parks, 2020; Neumaier et al., 2011]
- Build on recent derivative-based analysis

[Cartis, Fowkes & Shao, 2020]

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### Subspace DFO framework:

- Generate subspace of dimension  $p \ll n$  given by  $\operatorname{col}(Q_k)$  for random  $Q_k \in \mathbb{R}^{n \times p}$
- Build a low-dimensional model: find  $\hat{\boldsymbol{g}}_k \in \mathbb{R}^p$ ,  $\hat{H}_k \in \mathbb{R}^{p \times p}$  to get

$$f(\mathbf{x}_k + \mathbf{Q}_k \hat{\mathbf{s}}) \approx \hat{m}_k(\hat{\mathbf{s}}) = f(\mathbf{x}_k) + \hat{\mathbf{g}}_k^T \hat{\mathbf{s}} + \frac{1}{2} \hat{\mathbf{s}}^T \hat{H}_k \hat{\mathbf{s}},$$

- Solve subspace trust-region subproblem:  $\min_{\hat{s} \in \mathbb{R}^p} \hat{m}_k(\hat{s})$  s.t.  $\|\hat{s}\|_2 \leq \Delta_k$
- Benefits: fewer interpolation points needed, cheap linear algebra (everything in  $\mathbb{R}^p$ ).

# **Subspace DFO** — **Subspace Quality**

**Choice of subspace:** we need to make sure we search in 'good' subspaces (where there is potential to decrease f sufficiently).

The subspace at iteration k is well-aligned if

$$\|Q_k^T \nabla f(\mathbf{x}_k)\|_2 \ge \alpha \|\nabla f(\mathbf{x}_k)\|_2$$
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### **Key Assumption**

The subspace  $Q_k$  is well-aligned with probability  $1-\delta$  (whenever  $Q_k$  is resampled, independent of history), and  $\|Q_k\|_2 \leq Q_{\max}$ .

**Why?** If  $\|\nabla f(\mathbf{x}_k)\|_2 \geq \epsilon$ ,  $Q_k$  well-aligned and  $\hat{m}_k$  fully linear, then  $\|\hat{\mathbf{g}}_k\|_2 \geq \Omega(\epsilon)$ 

If there is still work to do, then the algorithm (probably) knows it

# **Subspace DFO Algorithm**

### RSDFO (Random Subspace DFO):

[model-based DFO, RSDFO-specific]

- 1. If FLAG, use previous  $Q_k = Q_{k-1}$  and construct fully linear subspace model  $\hat{m}_k$ .
- 2. Otherwise, generate random  $Q_k$  and construct subspace model  $\hat{m}_k$ .
- 3. If  $\|\hat{\boldsymbol{g}}_k\|_2$  small, ensure model fully linear and  $\Delta_k \sim \|\nabla f(\boldsymbol{x}_k)\|_2$ . [criticality]
- 4. Minimize model to get  $\mathbf{s}_k = Q_k \hat{\mathbf{s}}_k$ , evaluate  $f(\mathbf{x}_k + \mathbf{s}_k)$ .
- 5. Check sufficient decrease, then accept/reject step and update  $\Delta_k$ :
  - If decrease:  $x_{k+1} = x_k + s_k$  and  $\Delta_{k+1} = \gamma_{inc}\Delta_k$ , add  $x_{k+1}$  to model. [successful]
  - If no decrease and model not fully linear:  $x_{k+1} = x_k$  and  $\Delta_{k+1} = \Delta_k$ , make model fully linear. Set FLAG=TRUE. [model-improving]
  - If no decrease and model fully linear:  $x_{k+1} = x_k$  and  $\Delta_{k+1} = \gamma_{\text{dec}} \Delta_k$ . [unsuccessful]

# **Subspace DFO — Convergence**

### Theorem (Cartis & R., 2021)

If f is sufficiently smooth and bounded below,  $\gamma_{dec} > \gamma_{inc}^{-1/2}$  and  $\epsilon$  sufficiently small, then for some c, C > 0,

$$\mathbb{P}\left[ \mathsf{K}_{\epsilon} \leq \frac{\mathsf{C}}{\alpha^2 (1-\delta) \epsilon^2} \right] \geq 1 - \mathsf{e}^{-c\epsilon^{-2}},$$

where  $K_{\epsilon}$  is the first iteration with  $\|\nabla f(\mathbf{x}_k)\|_2 \leq \epsilon$ .

- ullet Matches usual  $\mathcal{O}(\epsilon^{-2})$  worst-case complexity bound with high probability
- ullet Implies  $\mathbb{E}\left[\mathcal{K}_{\epsilon}
  ight]=\mathcal{O}(\epsilon^{-2})$  and almost-sure convergence
- ullet Constant C depends on p (from fully linear error bounds), c depends on p and  $\delta$

# Convergence Proof — Sketch

**Proof sketch:** while  $\|\nabla f(\mathbf{x}_k)\|_2 > \epsilon$ , bound number of iterations across 6 cases.

### Good subspace:

- 1.  $\Delta_k$  large + successful: get  $f(\mathbf{x}_k) f(\mathbf{x}_{k+1}) \geq \Omega(\epsilon^2)$ , so happens  $\mathcal{O}(\epsilon^{-2})$  times.
- 2.  $\Delta_k$  large + unsuccessful: bounded by case #1 from  $\Delta_k$  management.
- 3.  $\Delta_k$  small + unsuccessful + good model: doesn't happen (Taylor accuracy)
- 4.  $\Delta_k$  small + successful: bounded by cases #3 and #5 from  $\Delta_k$  management
- 5.  $\Delta_k$  small + bad model: keep  $Q_k$  and  $\Delta_k$ , build good model (next time #3 or #4)

(extra difficulties: different  $\Delta_k$  large/small thresholds, 4  $\leftrightarrow$  5, criticality steps, ...)

### Bad subspace:

6. Happens with small probability  $\delta$ . Need  $\gamma_{\text{dec}} > \gamma_{\text{inc}}^{-1/2}$  to ensure  $\Delta_k$  not decreased too quickly in these iterations.

# Generating $Q_k$

For RSDFO to work, need to be able to generate  $Q_k$  such that

$$\|Q_k^T \nabla f(\mathbf{x}_k)\|_2 \ge \alpha \|\nabla f(\mathbf{x}_k)\|_2$$
 with probability  $\ge 1 - \delta$ .

If  $Q_k$  is a random orthonormal set (e.g. block coordinates), need  $p \sim \alpha n$ .

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Instead, make  $Q_k$  a Johnson-Lindenstrauss embedding, such as

- $Q_k$  has i.i.d. Gaussian entries  $\mathcal{N}(0,1/p)$
- $Q_k$  has s random nonzero entries per row, value  $\pm 1/\sqrt{s}$  with probability 1/2

Then, only need  $p \sim (1 - \alpha)^{-2} |\log \delta|$ , independent of n.

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$$\min_{\mathbf{x}\in\mathbb{R}^n} f(\mathbf{x}) = \frac{1}{2} \|\mathbf{r}(\mathbf{x})\|_2^2, \qquad \mathbf{r}(\mathbf{x}) \in \mathbb{R}^m$$

**Classical Gauss-Newton** 

**Derivative-Free Gauss-Newton** 

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#### **Classical Gauss-Newton**

#### **Derivative-Free Gauss-Newton**

• Linearize r at  $x_k$  using Jacobian

$$r(x_k+s) \approx m_k(s) = r(x_k) + J(x_k)s$$

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Jacobian not available: use

$$\boldsymbol{m}_k(\boldsymbol{s}) = \boldsymbol{r}(\boldsymbol{x}_k) + \boldsymbol{J}_k \boldsymbol{s}$$

• Find  $J_k$  using linear interpolation

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$$\boldsymbol{m}_k(\boldsymbol{s}) = \boldsymbol{r}(\boldsymbol{x}_k) + \boldsymbol{J}_k \boldsymbol{s}$$

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In both cases, get a local quadratic model

$$f(\boldsymbol{x}_k + \boldsymbol{s}) \approx m_k(\boldsymbol{s}) = \frac{1}{2} \|\boldsymbol{m}_k(\boldsymbol{s})\|_2^2$$

Implemented in state-of-the-art solver DFO-LS

[Cartis et al., 2019]

### **DFO for Least-Squares**

Standard method has first-order complexity  $\mathcal{O}(n^6\epsilon^{-2})$ : dependency on n between first & second order methods. [Cartis & R., 2019]

RSDFO with Gauss-Newton models gets dimension-independent  $\mathcal{O}(\epsilon^{-2})$  bound.

### **DFO for Least-Squares**

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RSDFO with Gauss-Newton models gets dimension-independent  $\mathcal{O}(\epsilon^{-2})$  bound.

#### **Practical considerations:**

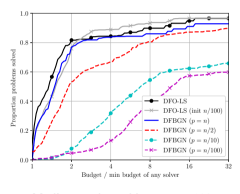
- Linear algebra cost of standard method is  $\mathcal{O}(mn^2 + n^3)$  flops per iteration from linear interpolation, RSDFO only needs  $\mathcal{O}(mp^2 + np^2)$
- Standard method reuses (possibly expensive) evaluations of r(x) across iterations, RSDFO has to resample all points from new subspace

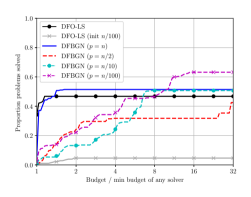
Key idea (DFBGN): use the locations of interpolation points to define the subspace  $\implies$  cheap linear algebra and fewer evaluations! If we have interpolation points  $\{x_k,y_1,\ldots,y_p\}$ , then make  $Q_k$  an orthonormal basis for  $\{y_1-x_k,\ldots,y_p-x_k\}$ .

# Numerical Results — low accuracy

DFBGN vs. DFO-LS (low accuracy  $au = 10^{-1}$ )

[% problems solved vs. # evals]





Medium-scale problems,  $n \approx 100$ 

Large problems  $n \approx 1000$ , 12hr timeout

DFBGN performance improves with larger p. Outperforms DFO-LS on large problems...

#### **Timeout Rate**

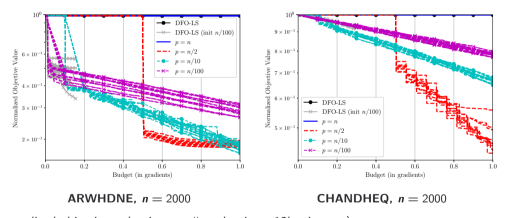
Proportion of large problems ( $n \approx 1000$ ) where solver times out (before usual termination):

Solver	Timeout
DFO-LS	93%
DFO-LS (init $n/100$ )	98%
DFBGN $(p = n/100)$	35%
DFBGN $(p = n/10)$	74%
DFBGN $(p = n/2)$	82%
$DFBGN\;(p=n)$	66%

#### ... because it doesn't time out

# Numerical Results — low budget

Other advantage: DFBGN progresses after  $p \ll n$  evaluations (important when n large)



(normalized objective reduction vs. # evaluations, 12hr timeout)

### **Conclusions & Future Work**

#### **Conclusions**

- Scalability of model-based DFO is currently limited (in theory & practice)
- New algorithms reduce linear algebra cost and iteration complexity
- Novel complexity analysis with dimension-independent bounds
- DFBGN outperforms state-of-the-art code on large-scale problems

#### **Future Work**

- Second-order complexity analysis
- Efficient implementation of subspace quadratic models
- Similar strategies for direct search DFO

[arXiv:2102.12016, Github: numerical algorithms group/dfbgn]

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### **Derivative-Free Block Gauss-Newton**

## Algorithm DFBGN (Derivative-Free Block Gauss-Newton):

- 1. Build low-dimensional model and calculate trust-region step  $oldsymbol{s}_k = Q_k \hat{oldsymbol{s}}_k$
- 2. Evaluate  $f(x_k + s_k)$ , accept/reject step, and update  $\Delta_k$  (as before)
- 3. Add  $x_k + s_k$  to interpolation set
- 4. Remove  $p_{drop} \ge 2$  points from the interpolation set
- 5. Add random orthogonal directions  $x_k + \Delta_k d$  until p+1 interpolation points

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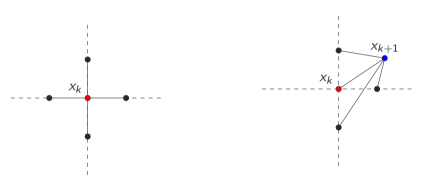
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#### **Comments:**

- $p_{drop} \ge 2$  ensures new direction(s) **d** added next iteration  $\implies Q_{k+1} \ne Q_k$ .
  - Practical choice:  $p_{\text{drop}}=2$  on success, p/10 otherwise (geometry-aware removal)
- Linear algebra cost  $\mathcal{O}(mp^2+np^2)$  vs. standard method  $\mathcal{O}(mn^2+n^3)$
- Package on Github: numerical algorithms group/dfbgn

# **General Objective Problems**

**General objective** case is much harder — rely on quadratic interpolation models.



2 points per subspace direction

After step, how to rotate subspace?

Subspace dimensions decoupled from interpolation directions  $\boldsymbol{y}_t - \boldsymbol{x}_k$