Sparse Grid Methods for Uncertainty Quantification

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- 1. Sparse grids
 - Construction principles and properties
 - Optimal sparse grids
 - Adaptive combination method
- 2. Application
 - Multi-scale viscoelastic flows

Motivation

- Numerical methods in uncertainty quantification:
 - Galerkin approach
 - Collocation technique
 - Discrete projection
- Needed on stochastic/parameter domain:
 - Approximation of integrals
 - Interpolation, especially for collocation
- Simple domains with product structure: $[-a,a]^d$, $I\!R^d$
- Issue: high- or even infinite-dimensional problems

Curse of dimension

- $f: \Omega^{(d)} \to I\!\!R$, $f \in V^{(r)}$, r isotropic smoothness
- Bellmann '61: curse of dimension M = #dof

$$\|f - f_M\|_{H^s} = C(d) \cdot M^{-r/d} |f|_{H^{s+r}} = O(M^{-r/d})$$

- Find situations where curse can be broken ?
- Trivial: restrict to r = O(d)

$$|f - f_M| = O(M^{-cd/d}) = O(M^{-c})$$

but practically not very relevant

 In any case: some smoothness changes with *d* or importance of coordinates decays successively (e.g. after suitable nonlinear transformation)

Sparse grid approach

- Basic principles:
 - 1-dim multilevel series expansion with proper decay
 - d-dim product construction
 - Trunctation of resulting multivariate expansion
- Effect:
 - reduction of cost complexity
 - nearly same accuracy as "full" product
 - necessary: certain smoothness requirements
 - adaptivity for detection of lower-dimensional manifolds

Simple example: Hierarchical basis



Tensor product hierarchical basis Generalization to higher dimension by tensor product



Table of subspaces $W_{l_1l_2}$



decay in x- and y-direction by 1/4 decay in diagonal direction by 1/16

Idea:

Omit points with small associated hierarchial coefficient values

Regular sparse grids

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Properties of regular sparse grids

Sparse grids Full grids $N \cong 2^n$ $O(N \log(N)^{d-1})$ instead of $O(N^d)$ Cost: L_2 -norm Accuracy: $O(N^{-2}\log(N)^{d-1})$ $O(N^{-2})$ $|\sum_{i=1}^{d} \frac{\partial^2 f}{\partial x_i^2}| \le c$ Smoothness: $\left|\frac{\partial^{2u}f}{\partial x^2}\right| \leq c$ Space, seminorm: H_{mix}^2 , $|f|_{2mix}$ H^2 , $|f|_2$

Mitigates the curse of dimension of conventional full grids Note: Higher regularity in mixed derivative, ~d

For wavelets, general stable multiscale systems: $O(N^{-2}(\log N)^{(d-1)/2})$

History of regular sparse grids

Re-invented several times:

- 1957 Korobov, Babenko
- 1963 Smolyak
- 1970 Gordon
- 1980 Delvos, Posdorf
- 1990 Zenger, G.
- 2000 Stromberg, deVore

2010 ????

hyperbolic cross points

blending method Boolean interpolation sparse grids hyperbolic wavelets

Application areas include:

- quadrature
- interpolation
- data compression

- solution of PDEs
- integral equations
- eigenvalue problems

Basic principles of sparse grids

- 1-dim multilevel sequence of operators and spaces $P_l: V^{(1)} \to V_l \qquad \qquad V_l \qquad \qquad l \in I\!\!N$
- Sequence of differences, telescopic approach

$$\Delta_l \coloneqq (P_l - P_{l-1}) : V^{(1)} \to V_l \ominus V_{l-1} \rightleftharpoons W_l$$

- d-dim. product construction $\mathbf{l} = (l_1, l_2, \dots, l_d) \in IN^d$ $\Delta_{\mathbf{l}} \coloneqq \bigotimes_{i=1}^d \Delta_{l_i} = \bigotimes_{i=1}^d (P_{l_i} - P_{l_i-1}) : V^{(d)} \to W_{\mathbf{l}} \qquad f_{\mathbf{l}} = \Delta_{\mathbf{l}}(f) \in W_{\mathbf{l}}$
- Appropriate truncation of resulting multivariate expansion $m^{d} \rightarrow \gamma = m^{d}$

$$\mathbb{N}^d \to \mathfrak{I} \subset \mathbb{N}^d$$

$$P = \sum_{\mathbf{l} \in N^d} \Delta_{\mathbf{l}} \quad \rightarrow \quad P_{\mathfrak{I}} = \sum_{\mathbf{l} \in \mathfrak{I}} \Delta_{\mathbf{l}}$$

Examples of multiscale expansions, 1d

- Integration: $P_l = Q_l : V^{(1)} \rightarrow V_l = IR$
 - Sequence of nested or non-nested point sets and weights, size: $n_l = l$ or $n_l = 2^l + 1$

=> various sparse grid quadrature rules

- Interpolation $P_l = I_l : V^{(1)} \to V_l$, approximation $P_l = A_l : V^{(1)} \to V_l$
 - Local piecewise polynomials, multiscale expansion: hierarchical basis, interpolets, wavelets, multilevel basis, size: $n_l = 2^l + 1$ $|W_l| = 2^{l-1}$

=> sparse grid finite element spaces

 Global polynomials: Fourier series, Chebyshev, Legendre, Hermite, Bernoulli polynomials

size
$$n_l = l$$
 or $n_l = 2^l + 1$ $|W_l| = 1$ or $|W_l| = 2^{l-1}$

=> total degree / hyperbolic cross approximation

Regular sparse grid approach

Index sets

$$\mathfrak{I}_{n}^{\text{full}} = \left\{ \mathbf{I} \in \mathbb{I} \mathbb{N}^{d} : |\mathbf{I}|_{\infty} = \max_{j=1,\dots,d} l_{j} \le n \right\}$$
$$\mathfrak{I}_{n}^{\text{sparse}} = \left\{ \mathbf{I} \in \mathbb{I} \mathbb{N}^{d} : |\mathbf{I}|_{1} = \sum_{j=1}^{d} l_{j} \le n + d - 1 \right\}$$

• The hierarchical representation is then

$$P_n^{\text{sparse}} = \sum_{|\mathbf{l}|_1 \le n+d-1} \Delta_{\mathbf{l}} \qquad P_n^{\text{sparse}}(f) = \sum_{|\mathbf{l}|_1 \le n+d-1} \Delta_{\mathbf{l}}(f)$$

- Other representations:
 - generating system
 - Lagrange system over SG points
 - semi-hierarchical
 - combination method

The combination technique

• A simple alternative representation is [G., Schneider, Zenger 91],

$$P_{n}^{\text{combi}} = \sum_{n \le |\mathbf{l}|_{1} \le n+d-1} (-1)^{n+d-|\mathbf{l}|_{1}-1} \begin{pmatrix} d-1\\ |\mathbf{l}|_{1} - 1 \end{pmatrix} P_{\mathbf{l}} \qquad P_{\mathbf{l}} := \bigotimes_{j=1}^{d} P_{l_{j}}$$

– Involves just the (anisotropic) full grid discretizations P_1 on different levels and linearly combines them



The combination technique

- Redundant representation but allows the simple reuse of existing code
- Completely parallel computation of the subproblems P_1
- Corresponds to a certain multivariate extrapolation method [Rüde 91]
- Necessary: Existence of a pointwise error expansion.
 - Euler-Maruyama of stochastic ODE: additive expansion (leading error term) of mean square error
- Multilevel-Monte Carlo is just 2-d combination method
 - Variance and bias for the two dimensions and a proper refinement rule which reflects the MC and the Euler-Maruyama rates [Gerstner12, Harbrecht, Peters, Siebenmorgen13]

A priori construction of sparse grids

- In general: Given
 - a class of functions and an error norm
 - an associated bound $b(\mathbf{l})$ for the benefit of $\Delta_{\mathbf{l}}$
 - a bound $c(\mathbf{l})$ for the cost of $\Delta_{\mathbf{l}}$
- We can a-priori derive a (quasi-) optimal sparse grid by solving a binary knapsack problem [Bungartz+G.03]

$$\max \sum_{\mathbf{l} \in N^d} \alpha_{\mathbf{l}} \cdot b(\mathbf{l}) \quad \text{such that} \quad \sum_{\mathbf{l} \in N^d} \alpha_{\mathbf{l}} \cdot c(\mathbf{l}) \leq C_{fix} \qquad \alpha_{\mathbf{l}} \in \{0, 1\}$$

and setting $\mathfrak{T}_C = \{ \mathbf{l} \in I N^d : \alpha_{\mathbf{l}} = 1 \}$

 Boils down to just sorting the quotients b(l)/c(l) of the benefit versus cost according to its size and taking the largest indices into account L^2 -norm-based sparse grids in H_{mix}^2

- Representation $f(\mathbf{x}) = \sum_{\mathbf{l}} f_{\mathbf{l}}(\mathbf{x})$ $f_{\mathbf{l}}(\mathbf{x}) \in W_{\mathbf{l}}$ $\mathbf{x} = (x_1, ..., x_d)$ $\mathbf{l} = (l_1, ..., l_d)$
- Cost per subspace $c(\mathbf{l}) = \dim(W_{\mathbf{l}}) = 2^{|\mathbf{l}-\mathbf{l}|_1}$
- Benefit for accuracy $\| f_1 \|_2 \le b(\mathbf{I}) = 3^{-d} \cdot 2^{-2|\mathbf{I}|_1} \cdot \| f \|_{\mathbf{2},mix} = O(2^{-2|\mathbf{I}|_1})$
- Choice of best subspaces ? Knapsack problem !
 => local benefit²/cost ratio

$$b^{2}(\mathbf{I})/c(\mathbf{I}) \approx \frac{2^{-4 \cdot |\mathbf{I}|_{1}}}{2^{|\mathbf{I}|_{1}}} = 2^{-5 \cdot |\mathbf{I}|_{1}} \qquad \Rightarrow \text{ regular sparse grid space}$$
$$V_{n}^{(d,opt)} = \bigoplus_{|\mathbf{I}|_{1}=n+d-1} W_{\mathbf{I}} \qquad \qquad l_{2} \qquad \qquad |\mathbf{I}|_{1}=n+d-1 \quad \text{isoline}$$
$$n \quad l_{1}$$

Anisotropic sparse grids

- Non-equal directions
 - Weighted Sobolev spaces [Sloan+Wozniakowski93]

$$H^{r}_{\gamma,mix}$$

- Anisotropic smoothness spaces [Gerstner+G. 98, G.+Zung15]

$$H_{mix}^{s_1,s_2,\ldots,s_d} = H^{s_1}(I_1) \otimes H^{s_2}(I_2) \otimes \cdots \otimes H^{s_d}(I_d)$$

- Different dimensions for different directions [G.+Harbrecht 11]

 $H^{s_1}(\Omega_1) \otimes H^{s_2}(\Omega_2) \otimes \ldots \otimes H^{s_d}(\Omega_d)$

- Via knapsack problem:
 - A priori construction of optimal anisotropic sparse index sets
 - log-terms disappear



Generalized sparse grids

- General index sets $\Im \subset \mathbb{N}^d$
- Downward closed set, no holes

 l_1

$$\mathbf{l} \in \mathfrak{I} \implies \mathbf{l} - e_j \in \mathfrak{I} \quad j = 1, \dots, d$$

- Associated sparse grid operator $P_{\Im} = \sum_{I \in \Im} \Delta_{I}$
- Associated space and associated function

$$V_{\mathfrak{I}} = \bigoplus_{\mathbf{l} \in \mathfrak{I}} W_{\mathbf{l}} \qquad P_{\mathfrak{I}} f = \sum_{\mathbf{l} \in \mathfrak{I}} \Delta_{\mathbf{l}}(f) = \sum_{\mathbf{l} \in \mathfrak{I}} f_{\mathbf{l}}$$

The combination technique

Can also be generalized to a given downward closed index set 3

$$P_{\mathfrak{I}} = \sum_{\mathbf{l} \in \mathfrak{I}} c_{\mathbf{l}} P_{\mathbf{l}}$$

Combination coefficient

$$c_{\mathbf{l}} = \sum_{\mathbf{z}=0}^{1} (-1)^{|\mathbf{z}|_{1}} \chi^{\Im} (\mathbf{l} + \mathbf{z})$$



with characteristic function $\chi^{\mathfrak{I}}$ on the index set \mathfrak{I}

- Again: just (anisotropic) full grid discretizations P₁ on different levels get linearly combined
- Note: many coefficients on the lower levels are zero

Tensor product sparse grids

- Examples:
 - space \times time, $d_1 = 3, d_2 = 1$, parabolic problems
 - space × parameters $d_1 = 3, d_2 = 10 20$

but smooth in parameter variables

- space × stochastics
$$d_1 = 3, d_2 = \infty$$

but analytic in stochastic variables

- Main result: Curse of dimension only w.r.t. the larger dimension and/or the lower smoothness [G.+Harbrecht11], [G.+Zung15]
- Time, parametrization and stochastic coordinates disappear in the overall complexity rate
 just space discretization matters

Sparse space-time grids

• Approximation error and necessary regularity [G.+Oeltz07]

$$\inf_{u_n \in V_n^0} \|u - u_n\|_{H^1(\Omega) \otimes L^2(0,T)} \le c \ 2^{-n} \|u\|_{H^2(\Omega) \otimes H^2((0,T))}$$

- Classical regularity theory (Ladyzenskaja, Wloka) $u \in H^2(\Omega) \otimes H^2((0,T))$
- Sparse space-time grids posses same approximation rate as full space-time grids but just cost complexity of space problem
- In each time slice there is a conventional full grid



Stochastic and parametric PDEs

Solutions of stochastic/parametric PDEs

 $\partial_t u(t, \mathbf{x}, \mathbf{y}) - \nabla \cdot A(\mathbf{x}, \mathbf{y}) \nabla f(t, \mathbf{x}, \mathbf{y}) = r(t, \mathbf{x}, \mathbf{y})$

live on product $(t, \mathbf{x}, \mathbf{y}) \in T \times \mathbf{X} \times \mathbf{Y}$

- of temporal domain T
- of spatial domain **X** with $d_1 = 1,2,3$
- and stochastic/parametric domain Y with d₂ large or even infinity.
- Often: Very high smoothness in y-part
 - Here: especially weighted analyticity for the different coordinates, decay in covariance [Cohen,Devore,Schwab10,11]
 - Then, even infinite-dimensional Y become treatable
- Sparse grid not only within stochastics but also between spatial, temporal and stochastic domain

Sparse grids and analytic functions

- Analytic regularity in polydisc with radii $\mathbf{r} := (r_1, \dots, r_d)$
- Sequence of smoothness indices $\mathbf{a} = (a_1, \dots, a_d) = \log(\mathbf{r})$
- With global polynomials: $|\Delta_{\mathbf{k}}(f)| \leq c \cdot e^{-(a_1k_1+\ldots+a_dk_d)}$
- Accuracy with respect to the involved #dof *M* [Beck,Nobile,Tamellini,Tempone12,14], [Tran,Webster,Zhang15], [G.+Oettershagen15]

$$gm(\mathbf{a}) = \left(\prod_{j=1}^{d} a_{j}\right)^{1/d} \qquad \kappa(d) = (d !)^{1/d} > d/e \qquad O(e^{-gm(\mathbf{a})\kappa(d)M^{1/d}}M^{(d-1)/d})$$

- For the infinite-dimensional case:
 - Logarithmic growth => algebraic rate [Todor,Schwab07], [Cohen,Devore,Schwab10,11]

$$\beta > 1$$
 $\sum_{j=1}^{\infty} \frac{1}{e^{a_j/\beta} - 1} < \infty$ $O(M^{-(\beta-1)})$ Stechkin's Lemma

 Linear growth => subexponential rate [G.+Oettershagen15], [Tran,Webster,Zhang15]
 3 [Tran,Webster,Zhang15]

$$\alpha > 0 \qquad a_j \ge \alpha \cdot j \qquad O(M^{-\frac{3}{8}\alpha \cdot \sqrt{\log(M)}} M^{1+\frac{\alpha}{4}} \log(M)^{-1/2})$$

Stechkin's Lemma can not show this rate but gives only an algebraic bound

Dimension-adapted sparse grids

- So far: function class known,
 - a-priori choice of best subspaces by optimization
 - size of benefit/cost ratio indicated if subspace is relevant => sparse grid patterns for \Im
- Now: for given single function *f*
 - adaptively build up a set $\ensuremath{\mathfrak{I}}$ of active indices
 - benefit $b(\mathbf{I}) := \|\Delta_{\mathbf{I}}(f)\|^2$, i.e. local error-indicator of f
 - cost $c(\mathbf{l}) = |W_1|$ for subspace W_1 ,
 - benefit/cost indicator $\varepsilon(\mathbf{l}) \coloneqq b(\mathbf{l})/c(\mathbf{l})$
 - refinement strategy to build new index set,
 - global stopping criterion => sparse grid pattern \mathfrak{J}
- Directions *T*×**X**×**Y** with product of different smoothness

The adaptive combination algorithm



Example

• Evolution of the algorithm:



- As any adaptive heuristics: may terminate too early
- If mixed regularity not present, refinement to the usual full grid

Application: Non-Newtonian fluids

- Classical Newtonian fluids: Obey Newton's law of viscosity, stress tensor is proportional to load/force
- But various complex fluids show strange behavior which is not correctly described







Barus effect

Weissenberg effect

tubeless siphon effect

Application: Non-Newtonian fluids

- Non-Newtonian fluids contain microstructures which are the reason for their unusual properties
 - Examples: paint, toothpaste, shampoo, blood, oils
- Polymeric fluids are a subset of non-Newtonian fluids
 - Long-chained molecules in a Newtonian solvent
 - Viscoelasticity due to interaction of elastic molecules and drag forces in basic flow
- A macroscopic model like the Navier Stokes equations
 + macrosopic extensions is no longer sufficient
- Needs to be augmented by model on the micro scale
 => Two scale modelling

Mathematical modelling

- The conservation equations for polymeric fluids are the same as for the Newtonian case, but the presence of polymer molecules contributes a polymeric extra-stress tensor $\boldsymbol{\tau}_{p}$ and an additional polymeric viscosity η_p such that the viscosity ratio $\beta < 1$
- The Navier-Stokes equations are now

$$\frac{\partial \mathbf{u}}{\partial t} + \mathbf{u} \cdot \nabla \mathbf{u} = \frac{1}{\text{Re}} \beta \Delta \mathbf{u} - \nabla p + \frac{1}{\text{Re}} \nabla \cdot \boldsymbol{\tau}_{p}$$
 conservation of momentum $\nabla \cdot \mathbf{u} = 0$

+ b.c., with Reynolds number Re and viscosity ratio $\beta = \frac{\eta_s}{1-1}$ $\eta_{s} + \eta_{n}$

 η_s solvent viscosity η_p polymeric viscosity

m

Microscopic modelling

 On the microsocopic scale, a polymer chain is modelled by a spring chain of K+1 beads



- Position **x** in physical space/flow domain $\Omega \subset \mathbb{R}^3$
- Orientations $\mathbf{q}_1, \dots, \mathbf{q}_K$ in configuration space $\Gamma \subset \mathbb{R}^{3K}$
- Probability to find chains at time *t* with position in $[\mathbf{x}, \mathbf{x} + d\mathbf{x}]$ and orientations in $[\mathbf{q}_1, \mathbf{q}_1 + d\mathbf{q}_1]...[\mathbf{q}_K, \mathbf{q}_K + d\mathbf{q}_K]$ $\psi: \Omega \times \Gamma \times [0,T] \rightarrow IR^+, \ (\mathbf{x}, \mathbf{q}_1, ..., \mathbf{q}_K, t) \rightarrow \psi \ (\mathbf{x}, \mathbf{q}_1, ..., \mathbf{q}_K, t)$

Fokker-Planck equation

- The function ψ is a pdf, i.e. $\psi \ge 0$, $\int_{\Gamma} \psi = 1$
- The application of Newton's 2nd law to the forces acting on chain leads to the Fokker-Planck equation $\frac{\partial \psi}{\partial t} + \nabla_{\mathbf{x}} \cdot (\mathbf{u} \psi) + \sum_{i=1}^{K} \nabla_{\mathbf{q}_{i}} \cdot \left((\nabla_{\mathbf{x}} \mathbf{u})^{T} \mathbf{q}_{i} \psi - \frac{1}{4De} \sum_{i=1}^{K} A_{ij} \mathbf{F}(\mathbf{q}_{i}) \psi \right) = \frac{1}{4De} \sum_{i=1}^{K} \sum_{j=1}^{K} A_{ij} \nabla_{\mathbf{q}_{i}} \cdot \nabla_{\mathbf{q}_{j}} \psi$

with Rouse matrix $A = [-1 \ 2 - 1]_{K}$

- Describes evolution of ψ under chain's spring forces
 F(q₁),..,F(q_K)
- Various models for spring force: Hooke: F(q) = q

FENE: $\mathbf{F}(\mathbf{q}) = \frac{\mathbf{q}}{1 - ||\mathbf{q}||^2 / b}, ||\mathbf{q}||^2 \le b, \text{ FENE-P: } \mathbf{F}(\mathbf{q}) = \frac{\mathbf{q}}{1 - \langle \mathbf{q}^2 \rangle / b}, \langle \mathbf{q}^2 \rangle \le b$

Coupling to the macro scale

- ψ represents polymeric configurations of micro-system
- Expectation in configuration space

$$\langle \cdot \rangle = \int_{\Gamma} \cdot \psi \ d \ \mathbf{q}_1 \dots d \ \mathbf{q}_K$$

• Coupling of internal configurations of micro system to macroscopic stress tensor via Kramer's expression $\boldsymbol{\tau}_p = C \sum_{i=1}^{K} \left(\langle \mathbf{q}_i \otimes \mathbf{F}(\mathbf{q}_i) \rangle - \mathbf{Id} \right)$

Constant C depends on model, Deborah number, viscosity ratio

- Issues with the Fokker-Planck equation
 - becomes more singular for higher values of *De* [Suli, Knezevic08]
 => extremely fine numerical resolution needed [Lozinski, Owen 03]
 3+3K = 3(K+1) dimensional + time-dependent => curse of dim.

Stochastic microscopic modelling

• There is a formal equivalence between the Fokker-Planck equation and stochastic partial differential eq.

$$d\vec{\mathbf{Q}}(\mathbf{x},t) = \left(-(\mathbf{u}\cdot\nabla)\vec{\mathbf{Q}}(\mathbf{x},t) + (\nabla\mathbf{u})\cdot\vec{\mathbf{Q}}(\mathbf{x},t) - \frac{1}{4De} A \mathbf{F}(\vec{Q}(\mathbf{x},t))\right) dt + \sqrt{\frac{1}{2De}} d\vec{\mathbf{U}}(t)$$

Deborah number $\rightarrow -\frac{1}{4De} A \mathbf{F}(\vec{Q}(\mathbf{x},t)) dt + \sqrt{\frac{1}{2De}} d\vec{\mathbf{U}}(t)$

- Describes evolution of *K* random fields $\vec{\mathbf{Q}} = (\mathbf{Q}_1, ..., \mathbf{Q}_K)^T$ that represent the configuration vector $\vec{\mathbf{q}} = (\mathbf{q}_1, ..., \mathbf{q}_K)^T$
- Brownian forces on the beads are modelled by the 3-dim. Wiener processes $W_i(t)$, i = 1,..., K+1
- The vector $\vec{\mathbf{U}}(t)$ consists of the component-wise differences

$$(\vec{\mathbf{U}}(t))_i = \mathbf{W}_{i+1}(t) - \mathbf{W}_i(t), \ i = 1,...,K$$

Stochastic microscopic simulation

- Brownian configuration fields (BCF) [Hulsen97] Random field $\overline{Q}(\mathbf{x},t)$ for configuration
- Discretization of x-space: the M_G grid cells make from the parabolic SPDE a system of SODEs (MoL)
- Discretization of SODE-system: Put M_B configuration fields in each of the M_G space grid cells and evolve their configuration discretely over time, i.e. all $M_G \cdot M_B$ configuration fields have fixed spatial positions (Eulerian view).



Stochastic microscopic simulation

- In each grid cell $k = 1, ..., M_G$ with center \mathbf{x}_k we solve/ integrate the stochastic DE for a number M_B of stochastic realizations $\bar{\mathbf{Q}}^{(j)}(\mathbf{x}_k, t), j = 1, ..., M_B$
- They are distributed according to the known equilibrium density ψ for t = 0
- But we do not know ψ for t > 0. Thus, we approximate the first moments $\langle \mathbf{Q}_i(\mathbf{x}_k, t) \otimes \mathbf{F}(\mathbf{Q}_i(\mathbf{x}_k, t)) \rangle$ in Kramer's relation as

$$\mathbf{T}_{p}(\mathbf{x}_{k},t) = C \sum_{i=1}^{K} \left(\left\langle \mathbf{Q}_{i}(\mathbf{x}_{k},t) \otimes \mathbf{F}(\mathbf{Q}_{i}(\mathbf{x}_{k},t)) \right\rangle - \mathbf{Id} \right)$$
$$\approx C \sum_{i=1}^{K} \left(\frac{1}{M_{B}} \sum_{j=1}^{M_{B}} \mathbf{Q}_{i}^{(j)}(\mathbf{x}_{k},t) \otimes \mathbf{F}(\mathbf{Q}_{i}^{(j)}(\mathbf{x}_{k},t)) - \mathbf{Id} \right)$$

i.e. we replace the integral by Monte Carlo quadrature

Numerics

- Navier Stokes equations:
 - Uniform grid cells, staggered grid, cell centers p, $\boldsymbol{\tau}_{p}$, cell faces \boldsymbol{u}
 - WENO for convective terms, 2nd order scheme for other terms
 - Euler or Crank-Nicolson in time, CFL-condition
 - Chorin-like projection method
- Microscale stochastic equations:
 - M_B stochastic samples for each grid cell => $M_G \cdot M_B$ samples
 - QUICK for convective terms
 - Explicit Euler-Maruyama, semi-implicit Euler for FENE
 - Same time step size as for NS equations
 - Variance reduction scheme with equilibrium control variates



Issues

- Code works as expected
- But: Huge memory requirements and huge computing times due to large numberM_B of realizations in each cell
- Example for 3D multi-scale problem
 - Flow domain Ω with
 - $M_G = 100 \times 100 \times 100$ grid cells
 - $M_B = 10.000$ stochastic realizations in each grid cell
 - Total memory requirements:
 - 8 MB for the pressure field *p*
 - 24 MB for the velocity field \boldsymbol{u}
 - 48 MB for the six independent components of $\boldsymbol{\tau}_p$
 - 75 GB*N for all the $M_G \cdot M_B$ stochastic variables
 - Some months of computing time





non-Newtonian



Sparse grid approach

- Consider our multiscale flow problem in more detail.
- We have the problem parameters: mesh width, time step size, stochastic realizations, springs
- How can we improve on computational complexity ?
 - Instead of MC use QMC
 - Multilevel-MC, MLQMC for stochastic ODEs (time + stoch.)

This is just a certain 2d combination technique/ sparse grid approach [Gerstner 12] [Harbrecht,Peters,Siebenmorgen13]

- Combination technique in all 3 discretization parameters
 i.e. for space x time x stochastics,
 and for model parameter K, i.e. x number of springs
- If the optimal combination formula is not a priori known: run the (dimension)-adaptive algorithm

Coordinates for the combination method



Indicators for the combination method

- Approximation of the vector ${f u}$ and the tensor ${f au}_p$
- Compute benefits $b(\mathbf{l})$ and costs $c(\mathbf{l})$ componentwise
- One index set for all components
- Weighted and scaled benefit/cost indicator

$$\varepsilon(\mathbf{l}) = \max\left\{\omega \cdot \frac{\|b(\mathbf{l})(\mathbf{u})\|_{2,2}}{c(\mathbf{l})(\mathbf{u}) \cdot \|b(1)(\mathbf{u})\|_{2,2}}, (1-\omega) \cdot \frac{\|b(\mathbf{l})(\tau_p)\|_{F,2}}{c(\mathbf{l}) \cdot \|b(1)(\tau_p)\|_{F,2}}\right\}$$

Scaling with initial level b(1) not necessary if $\omega = 0$ or $\omega = 1$

Example 1: Couette flow

- Non-Newtonian fluid in a 2D channel.
 - Fluid is at rest at initial time t = 0, De = 0.5
 - Shearing of fluid over time with rate $\dot{\gamma} = du / dy$
 - Linear spring force model (dumbbell, K=1)
 - Probability density function ψ : $(x, \mathbf{q}, t) \in \mathbb{R}^4 \rightarrow \psi$ $(x, \mathbf{q}, t) \in \mathbb{R}$

1d in space, 2d in configuration space and time-dependent



- Discretization:
 - Initial level $(1/\Delta x, 1/\Delta t, \text{ samples}) = (4, 16, 256)$
 - Refinement from level to level by factor *2
 - Error indicator $\omega = 1$, we are after error in **u**



Behaviour of adaptive combination technique



We asymptotically observe an anisotropic sparse grid structure



• Relative L_2 error of \mathbf{u}_1

- Comparison:
 - Full grid error
- $E(u_{6,6,6}) \approx 0.04$ $E(u_{7,7,7}) \approx 0.01$
- Cost (dof)

full grid

 $C(u_{6,6,6}) \approx 5.4 \times 10^8$ $C(u_{7,7,7}) \approx 4.3 \times 10^9$

sparse grid

 $C(u^{C}) \approx 4.6 \times 10^{7}$

Example 2: Steady extensional flow

- Non-Newtonian fluid in a 3D domain.
 - Steady uniaxial extensional flow, De = 1.0
 - Stress tensor $\boldsymbol{\tau}_p$ is aimed for
 - FENE force model, K-spring chain
 - We vary the number K of springs up to 5
 - Probability density function ψ : $(\mathbf{q},t) \in \mathbb{R}^{3K} \times \mathbb{R} \rightarrow \psi$ $(\mathbf{q},t) \in \mathbb{R}$

3N-dimensional in configuration, time-dependent, number of springs, no space

- Discretization
 - Initial level (samples, $1/\Delta t$, springs) = (1024, 2, 1)
 - Refinement for time and samples from level to level by factor *2, refinement for springs by +1
 - Error indicator $\omega = 0$, we are after error in $\boldsymbol{\tau}_p$



Example 2: Steady extensional flow



Behaviour of adaptive combination technique We observe:

- a sparse grid structure for all indices
- plus a nearly full grid
 between time and
 springs for the
 smallest sample size
- Different refinement:*2 versus +1
- Relative L_2 error for τ_{xx} of adaptive combination technique

Example 2: Steady extensional flow

• Convergence of model for rising number K of springs



- All results are computed on fine level with 2 million samples.
- Fixed stochastic time step width $\Delta t = 1/2048$

Concluding remarks

- Basic principles of sparse grids
- Optimization by knapsack problem
- Dimension-adaptive combination method
 - Solution of subproblems P_1 on levels I
 - Sparse grid approximation by linear combination
 - Refinement with hierarchical contributions $\Delta_{\mathbf{l}}~$ and local cost
- Application to non-Newtonian flow
 - Two-scale problem, stochastic microscale
- Adaptive combination method works on discretization directions (space x time x samples) and also for model parameters (... x springs)
- => Allows to couple discretization and modelling errors

The C library HCFFT G.+Hamaekers

- Hierarchical sparse grid interpolation based on:
 - Fast Fourier transform (FFT), fast Sine and Cosine transform
 - Fast Chebyshev transform, Fast Legendre transform
 - Various other polynomial transforms
- Different hierarchical bases for different dimensions
- Dyadic and arbitrary, non-dyadic refined grids
- Several types of general sparse grids
- Dimension-adaptive sparse grids
- For high precision: possible use of long double
- Freely available at

www.hcfft.org

The flow solver

 Code NAST3DGPF which is freely available at http://www.nast3dgpf.de/

