

QCD Anderson transition with overlap valence quarks and influence of external magnetic fields

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In collaboration with: L. von Smekal, A. D. M. Valois, G. Ramirez-Hidalgo

Based on: PRD 109 074512, PoS LATTICE2024 189, PoS LATTICE2025 123

- 1 ... with overlap valence quarks (on a twisted-mass sea)
- 2 ... and influence of external magnetic fields
- 3 Algebraic connections of overlap eigenmodes

Motivation

Fundamental transitions in QCD

- Chiral restoration
- Deconfinement

Open question

Is there a relation between both transitions?

- QCD Anderson transition **related to both phenomena**
- In this work: Focus on relation to chiral restoration

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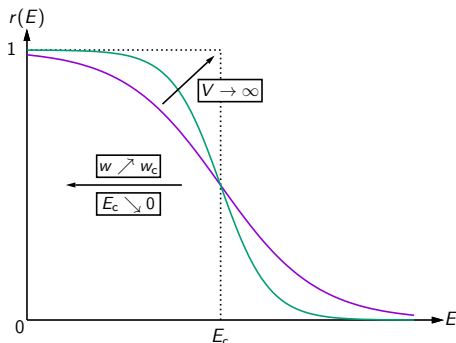
Anderson transition in condensed matter physics

[P. W. Anderson, PR **109** (1958) 1492], [F. Evers, A. D. Mirlin, RMP **80** (2008) 1355]

- Describes metal-insulator transition in disordered solids
- In metal phase **low-lying** eigenmodes of Hamiltonian **delocalized**
⇒ Conductivity
- Above critical disorder all eigenmodes **localized**
⇒ No conductivity

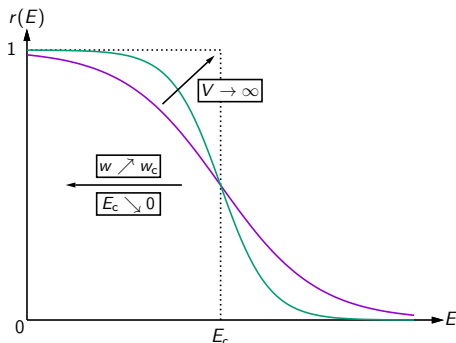
Anderson transition

- Delocalized modes separated from localized modes by energy threshold E_c (*mobility edge*)
- Above critical disorder strength w_c all modes localized



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Analogous transition in QCD

[M. Giordano, T. G. Kovács, Universe 7 (2021) 194]

- Hamilton operator
↳ *Dirac operator*
- Disorder strength
↳ *Temperature T*
- *Low-lying* modes *localized*
- Higher ones delocalized
- *Below T_0* all modes delocalized (no mobility edge)

... (de)confinement

- Eigenmodes tend to localize in sinks of Polyakov loop
[L. Holicki, E.-M. Ilgenfritz, L. von Smekal, PoS **LATTICE2018** (2019) 180]
- Quenched QCD: T_0 coincides with deconfining phase transition
[T. G. Kovács, R. Á. Vig, PRD **97** (2018) 014502]

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... chiral symmetry restoration/breaking

- Some studies suggest $T_0 = T_{pc}$ (pseudocritical temperature of chiral crossover, pion mass $m_\pi \neq 0$)
- No Goldstone bosons in chiral limit, if near-zero modes localized
[M. Giordano, JHEP **12** (2022) 103]
 $\Rightarrow T_0 \geq T_c$ (temperature of chiral phase transition, $m_\pi \rightarrow 0$)
- Near-zero modes produce chiral condensate (Banks-Casher relation)
[T. Banks, A. Casher, NPB **169** (1980) 103–125]

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Chiral lattice fermions

- Compute low-lying eigenmodes of overlap operator:

$$D_{\text{ov}} = \frac{\rho}{a} (1 + \text{sgn } K)$$

- Wilson kernel: $K = aD_W - \rho$
- Optimize locality with parameter $\rho \in (0, 2)$

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Gauge configurations

- From *twisted mass at finite temperature* (tmfT) collaboration
[F. Burger, E.-M. Ilgenfritz, M. P. Lombardo, A. Trunin, PRD **98** (2018) 094501]
 - Twisted mass Wilson fermions, Iwasaki gauge action
 - $N_f = 2 + 1 + 1$: two degenerate light, physical strange & charm quarks
 - Lattice spacing a from [N. Carrasco et al., NPB **887** (2014) 19–68]
 - m_π from [M. Werner et al., EPJA **56** (2020) 61]
 - T_{pc} from [A. Y. Kotov, M. P. Lombardo, A. Trunin, PLB **823** (2021) 136749]

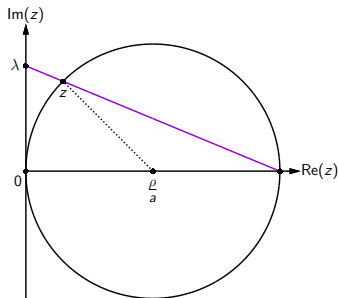
Observable and configurations

Localization measure

- Relative volume of eigenmode:

$$r(\lambda) = \frac{P^{-1}(\lambda)}{N_s^3 N_t} \in [0, 1]$$

- $P(\lambda) = \sum_n [v_\lambda(n)^\dagger v_\lambda(n)]^2$



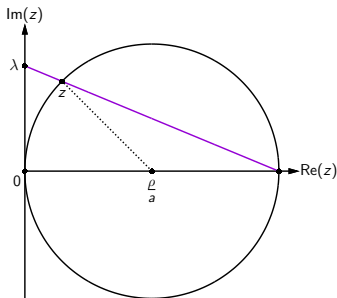
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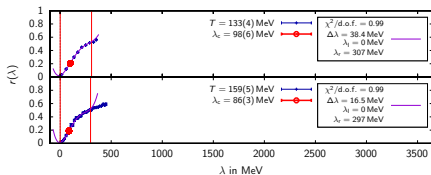
Set of ensembles	N_t	T/T_{pc}
D210 $N_s = 48$ $a = 0.0619(18)$ fm $L = 2.97(9)$ fm $m_\pi = 225(7)$ MeV $T_{pc} = 171(6)$ MeV	4	4.66(23)
	6	3.11(16)
	8	2.33(12)
	10	1.86(9)
	12	1.55(8)
	14	1.33(7)
	16	1.17(6)
	18	1.04(5)
	20	0.93(5)
	24	0.78(4)

- For $N_t = 18$:
 $T = 177(5)$ MeV $\approx T_{pc}$
- For $N_t = 24$:
 $T = 133(4)$ MeV $\approx T_c$

Relative volume for each T

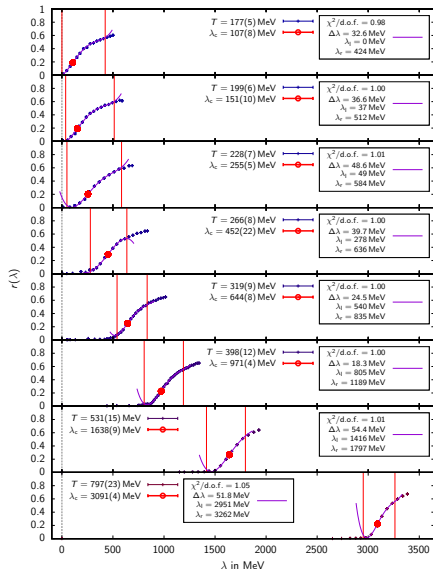
$$T_c = 132^{+3}_{-6} \text{ MeV}$$

[H.-T. Ding et al., PRL 123 (2019) 062002]



$$T_{pc} = 171(6) \text{ MeV}$$

- Inflection point λ_c as estimate for **mobility edge**
- **No expected vanishing** in the range $[T_c, T_{pc}]$



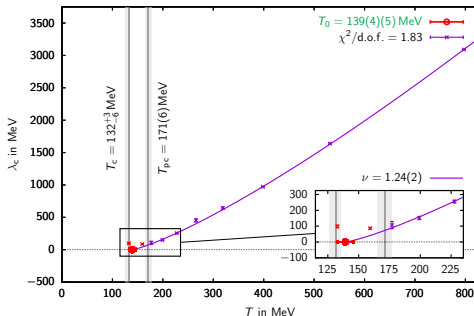
Temperature dependence

- Exclude **lowest two** temperatures
- λ_c seems to approach a **constant** here

- Zero **coincides with** T_c
 - $\nu \approx 1.44$ for unitary Anderson model
- [L. Ujfalusi, I. Varga, PRB **91** (2015) 184206]

Scaling fit

$$\lambda_c(T) = b(T - T_0)^\nu$$



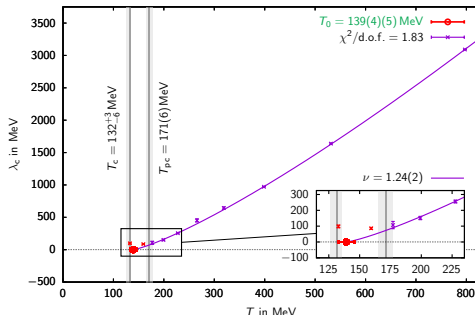
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Key question

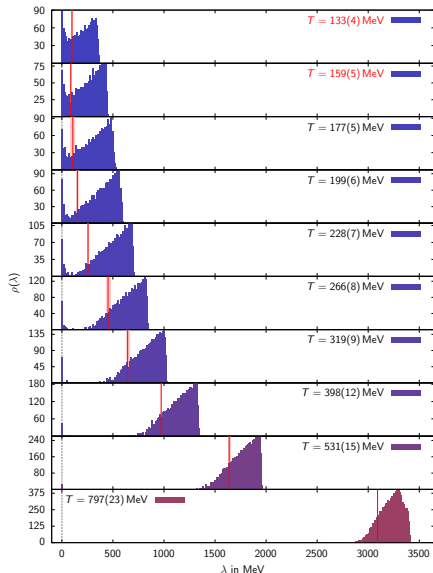
Why does the data below T_{pc} deviate from the fit?

Possible scenario

- If modes with low $r(\lambda)$ scale with $L^{d_{\text{IR}}}$, where $d_{\text{IR}} \in (0, 3]$
 $\Rightarrow$ Not localized
- Determine d_{IR} [A. Alexandru, I. Horváth, PRL 127 (2021) 052303]
 $\Rightarrow$ 2nd mobility edge $\lambda_{\text{IR}} = 0$
for $T > T_{\text{IR}} \in [200, 250]$ MeV
- Modes below λ_{IR} delocalized, higher ones localized

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 $\Rightarrow$ **2nd mobility edge $\lambda_{\text{IR}} = 0$** for $T > T_{\text{IR}} \in [200, 250]$ MeV
 - Modes below λ_{IR} delocalized, higher ones localized
-
- Decrease $T \Rightarrow \lambda_{\text{IR}}$ might increase and **annihilate λ_c**
 - Caution when λ_c hits infrared spectrum at $\approx T_{\text{pc}}$



- Investigate level spacing ratios:

$$\tilde{r}_n = \min \left(\frac{s_{n+1}}{s_n}, \frac{s_n}{s_{n+1}} \right) \in [0, 1]$$

[R. Shanker, H. Pandey, S. Sharma, JSPC
4 (2025) 100224]

- $s_n = \lambda_{n+1} - \lambda_n$
- $\langle \tilde{r} \rangle \approx 0.603$ for Gaussian unitary ensemble (**delocalized**)
- $\langle \tilde{r} \rangle \approx 0.386$ for Poisson distribution (**localized**)

Level spacing statistics

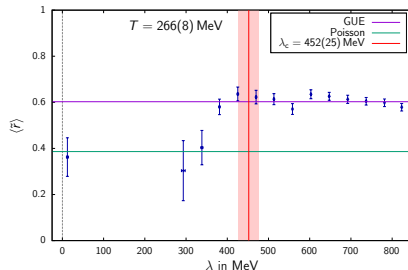
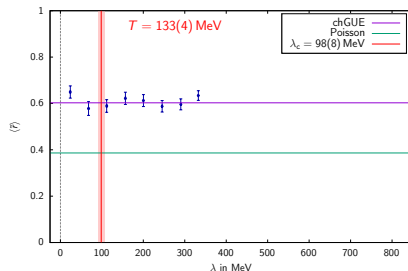
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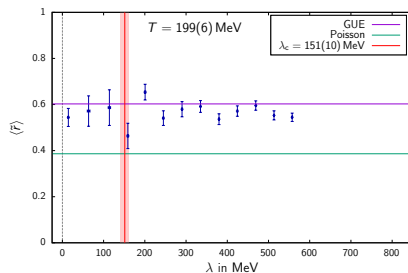
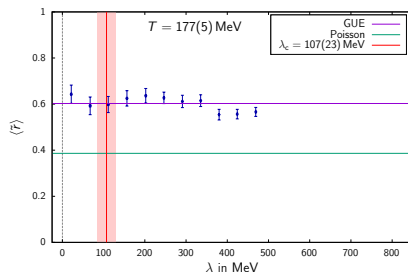
- According to $\langle \tilde{r} \rangle$ **no localization** at $T \approx T_c$ but for $T > T_{IR}$



Onset of localization?

- No localization at $T \approx T_{pc}$
- Slight indication of a dip at $T \approx 200$ MeV
- Seems to happen in the range $[T_{pc}, T_{IR}]$
- So far agreement with d_{IR} study
[X.-L. Meng et al., JHEP 12 (2024) 101]

- Improved statistics and smaller binsize required
- Larger volume & physical pion desirable (currently $L \approx 3$ fm, $m_\pi \approx 225$ MeV)
- Reduce computational costs



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Staggered fermions in magnetic fields

Staggered operator

$$D_{\text{st}}^f(n, l) = \sum_{\mu} \frac{\eta_{\mu}(n)}{2a} \left[U_{\mu}(n) u_{\mu}^f(n) \delta_{n+\hat{\mu}, l} - U_{\mu}^{\dagger}(n - \hat{\mu}) u_{\mu}^{f*}(n - \hat{\mu}) \delta_{n-\hat{\mu}, l} \right]$$

- Staggered phases:
 $\eta_1(n) = 1$, $\eta_{\mu}(n) = (-1)^{\sum_{\nu < \mu} n_{\nu}}$
- Flavor dependent $u_{\mu}^f(n) \in \text{U}(1)$
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Quantization condition on torus

- Magnetic flux ambiguous:
 $\Phi = q_f B S \quad \vee \quad \Phi_c = -q_f B (L^2 - S)$
- With $e^{i\Phi} \stackrel{!}{=} e^{i\Phi_c}$ and $q_d = -e/3$:
 $\Rightarrow \quad eB = 6\pi N_b / L^2, \quad N_b \in \mathbb{Z}$

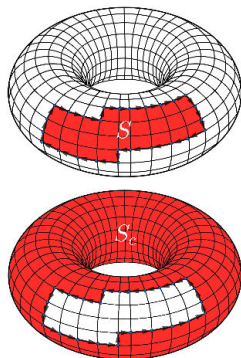


Figure: From A. D. M. Valois

Motivation

Magnetic fields **reduce** chiral & deconfinement transition temperature

[G. S. Bali et al., JHEP **02** (2012) 044]

⇒ Expect similar behavior for Anderson transition temperature T_0

(Inverse) magnetic catalysis

- Magnetic field enhances chiral condensate away from T_{pc}
- Condensate gets suppressed in region close to T_{pc} [F. Bruckmann, G. Endrődi, T. G. Kovács, JHEP **04** (2013) 112]

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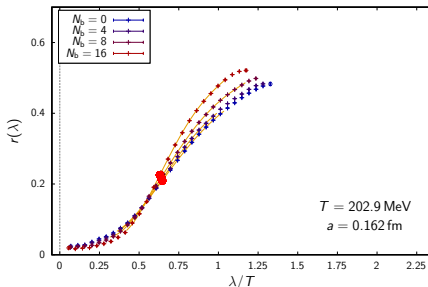
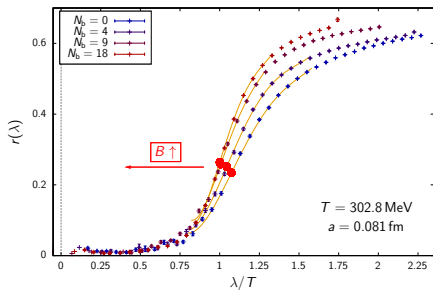
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Setup

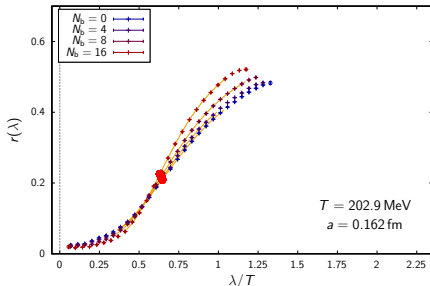
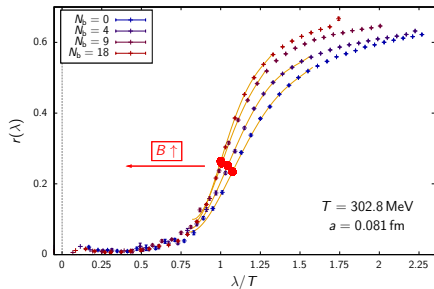
- Compute 400 eigenmodes of staggered operator
- 2 + 1 flavors of unimproved staggered fermions, tree-level Symanzik-improved gauge action
- $N_s = 24$, $N_t \in \{6, 8\}$
- Vary T via β

Relative volume: $N_s = 24$, $N_t = 6$



- B decreases λ_c at high T
- Seems to stay constant when Banks-Casher gap closes

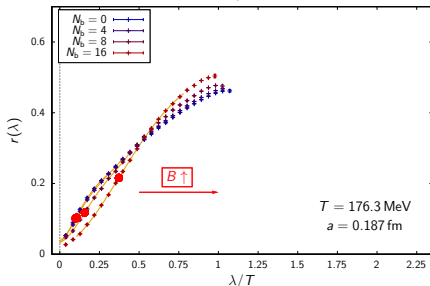
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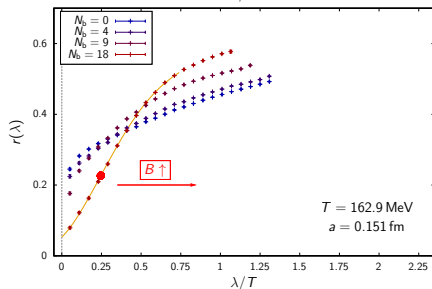
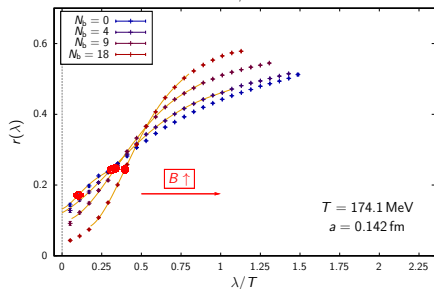
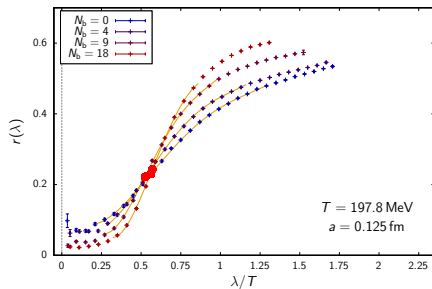
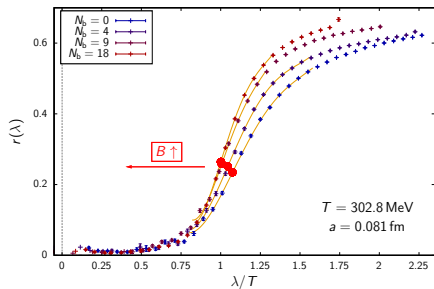
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Turn around

For lower T mobility edge gets increased by $B \Rightarrow T_0$ reduced



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Eigenpairs of D_{ov} from $D_{\text{ov}}^\dagger D_{\text{ov}}$

- $Dv = \lambda v \Leftrightarrow D(\gamma_5 v) = \lambda^*(\gamma_5 v)$
 $\Rightarrow$ Sufficient to compute just upper (or lower) semicircle
- Hermitian operators numerically more robust and cheaper
- γ_5 -hermiticity: $(\gamma_5 D)^\dagger = \gamma_5 D := H \Rightarrow D^\dagger D = H^2$ hermitian

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- v and $\gamma_5 v$ also eigenvectors of H^2 with degenerate eigenvalues $|\lambda|^2$
 - $[H^2, \gamma_5] = 0 \Rightarrow$ Can choose chiral basis $w_{\text{L/R}}$ of this eigenspace
 - Get rid of degeneracy by restricting to, e.g., right-handed subspace:
 $D^\dagger D = 2 + (1 + \gamma_5) \text{sgn } K$
 - Reconstruction:
 $\lambda_\pm = \frac{1}{2} \left(|\lambda|^2 \pm i|\lambda| \sqrt{4 - |\lambda|^2} \right) \Rightarrow v \propto (Hw_{\text{R}} - \lambda_\pm w_{\text{R}})$

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- However condition number gets worse: $\kappa(D^\dagger D) = \kappa(D)^2$

Eigenpairs of D_{ov} from $H = \gamma_5 D_{\text{ov}}$

- Condition number: $\kappa(\gamma_5 D) = \kappa(D)$
- $Hu_{\pm} = \pm|\lambda|u_{\pm}$ with **non**-chiral $u_{\pm}$ due to $[H, \gamma_5] \neq 0$
- But $w_{\text{L/R}} = P_{\text{L/R}} u_{\pm} \Rightarrow$ Formulas can still be applied
- Just positive (or negative) part of spectrum sufficient

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How to get rid of redundant part of spectrum?

- Simple chiral projection does not work here
- Reverse direction: $u_{\pm} \propto (v \pm \frac{\lambda}{|\lambda|} \gamma_5 v)$
- Idea: Filter initial vector u_0 for Krylov solvers
- Want something acting like $\Theta(H) = (1 + \text{sgn}(H))/2$
 - Approximate with Chebyshev polynomials
 - Amplify positive and suppress negative modes (e.g. also with Chebyshev polynomials)

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Thank you!

Implementation of the overlap operator

- High-dimensional matrix:

$$d = \underbrace{4}_{\text{Dirac}} \cdot \underbrace{3}_{\text{color}} \cdot \underbrace{\prod_{\mu=1}^4 N_{\mu}}_{\text{lattice sites}}$$

- $N_s = 24$, $N_t = 4$: $d \approx 660 \text{ k}$
 $\Rightarrow d^2 \approx 440 \text{ billion complex entries} \hat{\approx} 3.2 \text{ TB storage}$
(single precision)
- $N_s = 48$, $N_t = 24$: $\hat{\approx} 7.2 \text{ PB}$
- **No chance of storing the whole operator!**

- Implement matrix-vector product $D_{\text{ov}} v$ instead
- Rational approximation for inverse square root:

$$\begin{aligned} \text{sgn } K &= \frac{K}{\sqrt{(\gamma_5 K)^2}} \\ &\approx K \left(\alpha_0 + \sum_{j=1}^k \frac{\alpha_j}{(\gamma_5 K)^2 + \beta_j} \right) \end{aligned}$$

- Solve system of equations $\forall j$:

$$((\gamma_5 K)^2 + \beta_j) w_j = v$$

- Conjugate gradient method

Krylov subspace methods

- Due to high dimension **entire diagonalization not feasible!**
- Compute small part of spectrum using Krylov subspace methods

- p -dimensional Krylov subspace to matrix A and start vector x_1 :

$$\mathcal{K}_p(A, x_1) = \text{span}\{x_1, Ax_1, A^2x_1, \dots, A^{p-1}x_1\}$$

- Let $u_1, u_2, \dots, u_p$ be an orthonormal basis of the Krylov subspace
- Projection onto subspace:

$$H = U^\dagger AU$$

- Compute eigenvalues θ_i and eigenvectors y_i of H
- Approximate eigenpairs of A :
 $\lambda_i \approx \theta_i$ (Ritz values)
 $v_i \approx Uy_i$ (Ritz vectors)
- Best approximations in $\mathcal{K}_p(A, x_1)$

Arnoldi method

- Successively extend Krylov subspace with $\omega := Au_j$
- Compute matrix elements $H_{ij} = u_i^\dagger \omega$
- Orthogonalize ω to $u_1, u_2, \dots, u_j$
- Compute $H_{j+1,j} = \|\omega\|_2$ and set $u_{j+1} = \omega / \|\omega\|_2$

- H takes upper Hessenberg form of dimension p
- Arnoldi decomposition:

$$AU = UH + (Au_p)e_p^\dagger$$

- Krylov decomposition:

$$AU = UB + u_{p+1}b_{p+1}^\dagger$$

- **Krylov-Schur:** B has 1×1 or 2×2 blocks on the diagonal

Repeat

- Build Krylov decomposition using Arnoldi method
- Transform into Krylov-Schur decomposition
- Reorder blocks according to desired eigenvalues
- Truncate Krylov subspace