

# <sup>1</sup> KPM.jl: A Julia Package for Kernel Polynomial Method in Condensed Matter Physics

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## Software

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## <sup>11</sup> Summary

<sup>12</sup> The Kernel Polynomial Method (KPM) is a numerical technique for approximating spectral  
<sup>13</sup> and response functions of large Hermitian operators without full diagonalization ([Weisse et al., 2006](#)).<sup>14</sup> KPM expands spectral functions in Chebyshev polynomials and evaluates them  
via recursive matrix–vector multiplications, using stochastic trace estimation with random  
vectors to compute traces efficiently. Carefully chosen kernels (e.g., Jackson) suppress Gibbs  
oscillations and improve convergence of the truncated Chebyshev series. Because the dominant  
cost is sparse matrix–vector products, KPM scales (nearly) linearly with system size and can  
be applied to extremely large sparse Hamiltonians to compute quantities such as the density of  
states (DOS), local density of states (LDOS), and frequency-dependent response functions  
([João & Lopes, 2019](#)).

KPM.jl is a Julia implementation of KPM for tight-binding models in condensed matter physics,  
providing a high-performance, user-friendly toolbox. KPM.jl targets large sparse Hamiltonians,  
integrates with CUDA.jl for automatic GPU acceleration when available, and is designed  
for scalability and ease of integration into Julia-based workflows for large-scale spectral and  
transport calculations.

## <sup>27</sup> Statement of need

<sup>28</sup> KPM.jl provides a high-performance implementation of the Kernel Polynomial Method tailored  
<sup>29</sup> to tight-binding models. By avoiding full diagonalization and using Chebyshev expansions with  
<sup>30</sup> stochastic trace estimation, the package enables spectral and transport calculations on very  
<sup>31</sup> large sparse Hamiltonians that are infeasible with exact diagonalization (ED). It fills the gap  
<sup>32</sup> between low-level C/Fortran libraries and interactive, reproducible Julia workflows by offering:

- <sup>33</sup> 1. **Scalability:** Capable of handling large sparse matrices representing realistic tight-binding  
models.
- <sup>35</sup> 2. **Versatility:** Support for DOS and LDOS, as well as linear (DC conductivity) and nonlinear  
optical response functions.
- <sup>37</sup> 3. **Performance:** Automatic GPU acceleration via CUDA.jl when available.
- <sup>38</sup> 4. **Integration:** As a Julia package, it easily interfaces with other tools in the Julia ecosystem  
for Hamiltonian generation and data analysis.

<sup>40</sup> In condensed matter physics, understanding the effects of disorder, interactions, and complex  
<sup>41</sup> lattice geometries often requires numerical simulations of very large Hamiltonians. Traditional

42 ED methods for computing the full spectrum are limited to relatively small system sizes  
43 (typically  $N \sim 10^4$  states), which can be insufficient to resolve the spectral features of  
44 disordered systems or incommensurate structures like twisted bilayer graphene. KPM.jl is  
45 especially useful for studies of disorder, Moiré systems, and topological materials where large  
46 system sizes ( $N \sim 10^7$  states) are essential to capture realistic spectral and transport behavior.

## 47 Software design

48 KPM.jl is designed with modularity and performance in mind. The package operates on sparse  
49 Hamiltonians and computes Chebyshev moments using stochastic trace estimation with random  
50 vectors. The core functionality is divided into three main tiers corresponding to the complexity  
51 of the response function:

- 52   ■ **Density of States (1D):** The kpm\_1d function computes the Chebyshev moments for the  
53   DOS. It supports stochastic estimation using multiple random vectors (NR) and allows  
54   users to supply custom input vectors to compute the LDOS. The moments are converted  
55   to the spectral density  $\rho(E)$  using KPM.dos.
- 56   ■ **Linear Response (2D):** For transport properties like DC conductivity, kpm\_2d calculates  
57   the moments required for the Kubo-Greenwood formula involving two current operators  
58   ( $J_x, J_y$ ). The function d\_dc\_cond processes these moments to obtain the energy-  
59   dependent conductivity  $\sigma_{xy}(E)$ .
- 60   ■ **Nonlinear Response (3D):** The package includes specialized routines (kpm\_3d) for  
61   frequency-dependent nonlinear responses, such as the Circular Photogalvanic Effect  
62   (CPGE). This involves computing moments for three operators ( $J_x, J_y, J_z$ ) and post-  
63   processing them (d\_cpge) to extract the second-order conductivity  $\chi_{xyz}(\omega_1, \omega_2)$ .

64 The software automatically detects available hardware and offloads matrix-vector multiplications  
65 to a GPU if a compatible CUDA device is present (KPM.whichcore()), ensuring efficient  
66 performance on modern clusters.

## 67 Research impact statement

68 KPM.jl has been utilized in numerous studies to investigate electronic structure and response  
69 in complex materials:

- 70   1. **Moiré systems and disorder:** Enabled large-scale simulations of twisted bilayer graphene  
71   and demonstrated how twist-angle disorder broadens flat bands and modifies spectral  
72   features (Chang et al., 2024; Fu et al., 2020; Wilson et al., 2020; Yi et al., 2022).
- 73   2. **Experimentally relevant transport calculations:** Applied to transport modeling based on  
74   tight-binding models derived from DFT+U to directly compare theory with experiment  
75   (Liu et al., 2021).
- 76   3. **Quasiperiodic potentials for topological material:** Used to compute conductivity and  
77   disorder-averaged spectral properties and Green's function in disordered materials (Fu et  
78   al., 2021; Yi et al., 2022).
- 79   4. **Nonlinear optical responses and disordered topological material:** Employed to calculate  
80   nonlinear responses such as the CPGE in disordered topological semimetals and to  
81   identify spectral signatures of surface and bulk phase transitions in disordered axion  
82   insulators (Grindall et al., 2025; Wu et al., 2024).
- 83   5. **Method benchmarking and numerical validation:** Served as a reference implementation for  
84   comparing and validating new Chebyshev-regularization and spectral-density algorithms  
85   (Yi et al., 2025).

86 These applications demonstrate the package's capability to tackle cutting-edge problems in  
87 condensed matter theory.

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