

HPDDM

<https://github.com/hpddm/hpddm>

All keywords must be prefixed by `-hpddm_`. If a value is specified in the column “Default”, this value is used when the corresponding option is not set by the user. When no default value is specified but the corresponding option is set by the user, the option is true (represented internally by 1). If the option is not set, its value is false (represented internally by 0). Options highlighted in **red** should be reserved to expert users.

Keyword	Description	Possible values	Default
<code>help</code>	Display available options	Anything	
<code>version</code>	Display information about HPDDM	Anything	
<code>config_file</code>	Load options from a file saved on disk	String	
<code>tol</code>	Relative decrease in residual norm to reach in order to stop iterative methods	Numeric	10^{-6}
<code>max_it</code>	Maximum number of iterations of iterative methods	Integer	100
<code>verbosity</code>	Level of output (higher means more displayed information)	Integer	
<code>compute_residual</code>	Print the residual after convergence	12, 11, 1infy	
<code>push_prefix</code>	Prepend a prefix for all following options (use <code>-hpddm_pop_prefix</code> when done)		
<code>reuse_preconditioner</code>	Do not factorize again the local matrices when solving subsequent systems	Boolean	
<code>local_operator_spd</code>	Assume the local operator is symmetric positive definite	Boolean	
<code>orthogonalization</code>	Method used to orthogonalize a vector against an orthogonal basis	cgs, mgs	cgs
<code>dump_matri(ces x_[[:digit:]]+)</code>	Save either one or all local matrices to disk	String	
<code>dump_eigenvectors(_[[:digit:]]+)?</code>	Save either one or all local eigenvectors to disk	String	
<code>krylov_method</code>	Type of iterative method used to solve linear systems	gmres, bgmres, cg, bcg, gcrodr, bgcrodr, bfbcg, richardson, none	gmres
<code>enlarge_krylov_subspace</code>	Split the initial right-hand side into multiple vectors	Integer	1
<code>gmres_restart</code>	Maximum number of Arnoldi vectors generated per cycle	Integer	40
<code>variant</code>	Left, right, or variable preconditioning	left, right, flexible	right
<code>qr</code>	Method used to perform distributed QR factorizations	cholqr, cgs, mgs	cholqr
<code>deflation_tol</code>	Tolerance when deflating right-hand sides inside block methods	Numeric	
<code>recycle</code>	Number of harmonic Ritz vectors to compute	Integer	
<code>recycle_same_system</code>	Assume the system is the same as the one for which Ritz vectors have been computed	Boolean	
<code>recycle_strategy</code>	Generalized eigenvalue problem to solve for recycling	A, B	A
<code>recycle_target</code>	Criterion to select harmonic Ritz vectors	SM, LM, SR, LR, SI, LI	SM
<code>richardson_damping_factor</code>	Damping factor using in Richardson iterations	Numeric	1.0
<code>eigsolver_tol</code>	Tolerance for computing eigenvectors by ARPACK or LAPACK	Numeric	10^{-6}
<code>geneo_nu</code>	Number of local eigenvectors to compute for adaptive methods	Integer	20
<code>geneo_threshold</code>	Threshold for selecting local eigenvectors for adaptive methods	Numeric	
<code>geneo_estimate_nu</code>	Estimate the number of eigenvalues below a threshold using the inertia of the stencil	Numeric	
<code>geneo_force_uniformity</code>	Ensure that the number of local eigenvectors is the same for all subdomains	min, max	
<code>master_p</code>	Number of master processes	Integer	1
<code>master_distribution</code>	Distribution of coarse right-hand sides and solution vectors	centralized, sol	centralized
<code>master_topology</code>	Distribution of the master processes	0, 1, 2	0
<code>master_assembly_hierarchy</code>	Hierarchy used for the assembly of the coarse operator	Integer	
<code>master_aggregate_size</code>	Number of master processes per MPI sub-communicators	Integer	master_p

master_dump_matrix	Save the coarse operator to disk	String	
master_exclude	Exclude the master processes from the domain decomposition	Boolean	
master_spd	Assume the coarse operator is symmetric positive definite	Boolean	

When using Schwarz methods, there are additional options.

Keyword	Description	Possible values	Default
schwarz_method	Type of Schwarz preconditioner used to solve linear systems	ras, oras, soras, asm, osm, none	ras
schwarz_coarse_correction	Type of coarse correction used in two-level methods	deflated, additive, balanced	

When using substructuring methods, there is an additional option.

Keyword	Description	Possible values	Default
substructuring_scaling	Scaling used in the definition of the Schur complement preconditioner	multiplicity, stiffness, coefficient	multiplicity

When using MKL PARDISO as a subdomain solver (resp. coarse operator solver), there are additional options, cf. <https://software.intel.com/en-us/node/470298> (resp. <https://software.intel.com/en-us/node/590089>).

Keyword	Description	Possible values
mkl_pardiso_iparm_(2 8 1[013] 2[147])	Integer control parameters of MKL PARDISO for the subdomain solvers	Integer
master_mkl_pardiso_iparm_(2 1[013] 2[17])	Integer control parameters of MKL PARDISO for the coarse operator solver	Integer

When using MUMPS as a subdomain solver (resp. coarse operator solver), there are additional options, cf. <http://mumps.enseeiht.fr/index.php?page=doc>.

Keyword	Description	Possible values
mumps_icntl_([678] 1[234] 2[3789] 35)	Integer control parameters of MUMPS for the subdomain solvers	Integer
mumps_cntl_([123457])	Real control parameters of MUMPS for the subdomain solvers	Numeric
master_mumps_icntl_([678] 1[234] 2[3789] 35)	Integer control parameters of MUMPS for the coarse operator solver	Integer
master_mumps_cntl_([123457])	Real control parameters of MUMPS for the coarse operator solver	Numeric

When using *hypre* as a coarse operator solver, there are additional options, cf. <http://acts.nersc.gov/hypre/#Documentation>.

Keyword	Description	Possible values	Default
master_hypre_solver	Solver used by <i>hypre</i> to solve coarse linear systems	fgmres, pcg, amg	fgmres
master_hypre_tol	Relative convergence tolerance	Numeric	10^{-12}
master_hypre_max_it	Maximum number of iterations	Integer	500
master_hypre_gmres_restart	Maximum number of Arnoldi vectors generated per cycle when using FlexGMRES	Integer	100
master_boomeramg_num_sweeps	Number of sweeps	Integer	1
master_boomeramg_max_levels	Maximum number of multigrid levels	Integer	10
master_boomeramg_coarsen_type	Parallel coarsening algorithm	Integer	6
master_boomeramg_relax_type	Smoother	Integer	3
master_boomeramg_interp_type	Parallel interpolation operator	Integer	0

When using ARPACK as an eigensolver, there is an additional option.

Keyword	Description	Possible values
arpack_ncv	Number of Lanczos basis vectors generated in one iteration	Integer