	Туре	Description	Mandatory
f ormat	string	"ES-HDF"	yes
version	int[3]	major,minor,patch	yes
creator supercell			
	double[3][3]	primitive lattice vectors	yes
boundary_conditions	int[3]	1 for periodic, 0 for open	yes
atoms mumber of atoms	int	number of atoms in primitive cell	yes yes
mumber_of_species	int	number of atomic species	yes
positions	double[N][3] double[N][3]	absolute positions in bohr position in units of primitive vectors	yes
reduced_positions species ids	int[N]	Indices of species listed below	no yes
species_0		·	•
atomic_number mass	int double	atomic number of the species atomic mass	yes
name	int	name of atomic species (e.g. C)	yes yes
m pseudopotential	int	pseudopotential file name	no
	int	charge of pseudo-ion	yes
i species_1			
electrons	ton		yes
<pre>number_of_electrons number_of_kpoints</pre>	int int	number of electrons in the primitive of number of k-points	cell yes yes
I number_of_spins	int	number of spins (1 or 2)	yes
functional total energy	string	exchange-correlation functional	no
total_energy psi_r_is_complex	int int	total energy in Hartrees 1 if real-space orbitals are complex	yes yes
psi_r_mesh	int[3]	number of points in real-space mesh	yes
inumber_of_atomic_orbitals	int int	number of atomic orbitals (may be 0)	
■ have_dpsi □ kpoint 0	IIIL	1 if derivatives of orbitals are present	t no yes
number_of_gvectors	int	number of reciprocal lattice vectors	yes
gvectors reduced k	int[N][3] double[3]	g-vectors in units of primitive recipro k-point in units of primitive reciprocla	
weight	double	weight for this k-point	ı lattice yes yes
spin_0			VAS
number_of_states ccupations	int double[N]	number of orbitals for this k-point and occupation for each state	d spin yes
eigenvalues	double[N]	eigenvalues in Hartrees	yes yes
state_0			yes
	double[][2]	complex coefficients corresponding to	
	double[N_x][N_y][N_z][1 or 2] double[N_{sp}][$(I_{max}+1)^2$][2]	real or complex coefficients on psi_r_ radial functions for atomic orbital rep	
radial_spline_1	double[N_{sp}][$(I_{max}+1)^2$][2]	radial functions for atomic orbital rep	
: = poly_coefs_0	$double[N_{sp}][(I_{max}+1)^2][2]$	polynomial coefficients for atomic or	
poly_coefs_1	$double[N_{sp}][(I_{max}+1)^2][2]$	polynomial coefficients for atomic orl	oital rep. no
L C state_1			
spin_1			
kpoint_1			
atomic_orbital_0			no
<u> </u>	int	maximum I channel for Y ^I expansion	n yes
	double int	radius of muffin tin for atomic orbital order for polynomial expansion near	yes origin yes
polynomial_radius	double	radius below which to use polynomia	
spline_radius	double	maximum radius for spline data	yes
spinic_points	int	number of spline data points	yes
atomic_orbital_1			
density			no
<pre> number_of_gvectors gvectors</pre>	int int[N][3]	number of gvectors for the density g-vectors in units of primitive recipro	yes cal lattice yes
gvectors mesh	int[3]	number of points in real-space mesh	cal lattice yes yes
= spin_0			
density_r density_g	double[N _x][N _y][N _z] double[][2]	real coefficients on mesh complex coefficients corresponding to	yes o g-vectors yes
spin_1	4048/C[][2]	complex coefficients corresponding to	og vectors yes
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