

# Species file format reference

version lithium

# About this Document

This document describes the file format for the species definitions.

# Part I

# Input Elements

# 1 Element: spdb

Species-database element contains the species element sp

Contains: sp XPath: /spdb

# 2 Element: sp

A species is an atom type definition containing all information to construct the basis functions.

Contains: muffinTin (1 times)

atomicState (1 times or more)

basis (1 times)
lorb (zero or more)

XPath: /spdb/sp

This element allows for specification of the following attributes:

```
chemicalSymbol (required), mass (required), z (required), name
```

#### 2.1 Attribute: chemicalSymbol

Chemical Symbol.

Type: ID Use: required

XPath: /spdb/sp/@chemicalSymbol

#### 2.2 Attribute: mass

Mass in  $m_e$ .

**Type:** fortrandouble (9.1)

Use: required

XPath: /spdb/sp/@mass

## 2.3 Attribute: name

Optional element name.

Type: stringUse: optional

XPath: /spdb/sp/@name

## 2.4 Attribute: z

Atomic number.

**Type:** fortrandouble (9.1)

Use: required XPath: /spdb/sp/@z

# 3 Element: muffinTin

This element gives the size of the muffin tin radius and the resolution of the radial functions.

**Type:** no content

XPath: /spdb/sp/muffinTin

This element allows for specification of the following attributes:

```
{\tt radialmeshPoints} \ (\overline{required}), \ radius \ (\overline{required}), \ rinf \ (\overline{required}), \\ rmin \ (\overline{required})
```

#### 3.1 Attribute: radialmeshPoints

Number of data points for radial atomic functions.

Type: integer Use: required

XPath: /spdb/sp/muffinTin/@radialmeshPoints

#### 3.2 Attribute: radius

The radius of the muffin tin sphere.

**Type:** fortrandouble (9.1)

Use: required

XPath: /spdb/sp/muffinTin/@radius

#### 3.3 Attribute: rinf

Radius from which the influence on the potential is regarded to be negligible.

**Type:** fortrandouble (9.1)

Use: required

XPath: /spdb/sp/muffinTin/@rinf

#### 3.4 Attribute: rmin

The radius where radial mesh begins.

**Type:** fortrandouble (9.1)

Use: required

XPath: /spdb/sp/muffinTin/@rmin

# 4 Element: atomicState

The atomicState element lists the atomic states that schould be used to approximate the wavefunction in the sphere. They can be marked as core or none core electrons by the core attribute. Core electrons are threated separately by numeric itegration.

**Type:** no content

XPath: /spdb/sp/atomicState

This element allows for specification of the following attributes:

```
core (required), kappa (required), 1 (required), n (required),
occ (required)
```

#### 4.1 Attribute: core

If true, state is threated as core state in the calculation.

Type: boolean Use: required

XPath: /spdb/sp/atomicState/@core

# 4.2 Attribute: kappa

Relativistic quantum number.

Type: integer Use: required

XPath: /spdb/sp/atomicState/@kappa

#### 4.3 Attribute: 1

Azimuthal quantum number.

Type: integer Use: required

XPath: /spdb/sp/atomicState/@l

#### 4.4 Attribute: n

Principal quantum number.

Type: integer Use: required

XPath: /spdb/sp/atomicState/@n

## 4.5 Attribute: occ

Ocuppation number.

**Type:** fortrandouble (9.1)

Use: required

XPath: /spdb/sp/atomicState/@occ

## 5 Element: basis

Defines APW basis.

Contains: wf (1 times or more)

exception (zero or more)

XPath: /spdb/sp/basis

This element allows for specification of the following attributes:

order (required)

#### 5.1 Attribute: order

Type: integer Use: required

XPath: /spdb/sp/basis/@order

# 6 Element: exception

This element allows for defining exceptions to the APW basis functions.

Contains: wf (1 times or more)

XPath: /spdb/sp/basis/exception

This element allows for specification of the following attributes:

1

#### 6.1 Attribute: 1

Spezifies the azimuthal quantum number for which the exception applies.

Type: integer Use: optional

XPath: /spdb/sp/basis/exception/@l

## 7 Element: lorb

Local orbital (APW+lo or LAPW+lo).

Contains: wf (1 times or more)
XPath: /spdb/sp/lorb

This element allows for specification of the following attributes:

1 (required)

#### 7.1 Attribute: 1

Azimuthal quantum number for which the local orbital is defined.

XPath: /spdb/sp/lorb/@l

# Part II

# Reused Elements

The following elements can occur more than once in the input file. Therefore they are listed separately.

# 8 Element: wf

Defines the radial part of an atomic wavefunction. This functions used to construct a lapse orbital. The actual basis functions for the calculation inside the MT are linear combinations of these and  $Y_{lm}$ .

Type: no content XPath: ./wf

Parent: /spdb/sp/basis

/spdb/sp/basis/exception

/spdb/sp/lorb

This element allows for specification of the following attributes:

matchingOrder (required), searchE (required), trialEnergy (required)

# 8.1 Attribute: matchingOrder

Gives the order of the derivative that must be matched to the plain wave.

Type: integer Use: required

XPath: ./wf/@matchingOrder

#### 8.2 Attribute: searchE

If true the energy of the radial wave function,  $E_0$  is optimized to match the boundary condition

$$\psi(R_{MT}) = 0. (1)$$

Type: boolean Use: required

XPath: ./wf/@searchE

# 8.3 Attribute: trialEnergy

Energy level of the radial wave function (initial condition for numerical radial Schrodinger equation)

**Type:** fortrandouble (9.1)

Use: required

XPath: ./wf/@trialEnergy

# 9 Data Types

The Input definition uses derived data types. These are described here.

## 9.1 Type fortrandouble

The type fortrandouble allows to use the letters "eEdDqQ" for exponent operators. This alters in what precision the number is parsed.