

Open access · Journal Article · DOI:10.1007/S00214-005-0655-Y

### Pseudopotentials for H to Kr optimized for gradient-corrected exchange-correlation functionals — Source link

Matthias Krack

Institutions: ETH Zurich

Published on: 25 May 2005 - Theoretical Chemistry Accounts (Springer-Verlag)

#### Related papers:

- Separable dual-space Gaussian pseudopotentials
- Generalized Gradient Approximation Made Simple
- QUICKSTEP: Fast and accurate density functional calculations using a mixed Gaussian and plane waves
   approach
- · Relativistic separable dual-space Gaussian pseudopotentials from H to Rn
- · Gaussian basis sets for accurate calculations on molecular systems in gas and condensed phases.



## **ETH** zürich

# Pseudopotentials for H to Kr optimized for gradient-corrected exchange-correlation functionals

**Journal Article** 

Author(s): Krack, M.

Publication date: 2005-09

Permanent link: https://doi.org/10.3929/ethz-b-000031572

Rights / license: In Copyright - Non-Commercial Use Permitted

**Originally published in:** Theoretical Chemistry Accounts 114(1-3), <u>https://doi.org/10.1007/s00214-005-0655-y</u>

#### **REGULAR ARTICLE**

#### M. Krack

## **Pseudopotentials for H to Kr optimized for gradient-corrected exchange-correlation functionals**

Received: 25 November 2004 / Accepted: 18 December 2004 / Published online: 25 May 2005 © Springer-Verlag 2005

Abstract Pseudopotential parameter sets for the elements from H to Kr using the relativistic, norm-conserving, separable, dual-space Gaussian-type pseudopotentials of Goedecker, Teter, and Hutter (GTH) are presented as optimized for the gradient-corrected exchange-correlation functionals of Becke, Lee, Yang, and Parr (BLYP), Becke and Perdew (BP), and Perdew, Burke, and Ernzerhof (PBE). The accuracy and reliability of the GTH pseudopotentials is shown by calculations for a series of small molecules.

Keywords Goedecker pseudopotentials  $\cdot$  Gaussian plane waves method  $\cdot$  Quickstep  $\cdot$  CP2K  $\cdot$  Gradient corrected functionals

#### **1** Introduction

The development of new electronic structure methods accompanied by the rapid development of the computer hardware during the last decades is causing a growing interest to use such methods for the investigation of complex chemical processes. However, the chemical processes of interest often involve heavy elements, in particular transition metals, as a key ingredient. A further complication is possibly introduced by the system size, since large model systems have to be employed in order to take environmental effects properly into account. An all-electron description of all atoms is computationally expensive and often not needed. Moreover, the inclusion of relativistic effects is indispensable for the heavy elements. The employment of effective pseudopotentials is known to be a valid method to reduce the computational effort in such cases, since the chemically inert core electrons are not explicitly considered in the calculation. Furthermore,

Contribution to Karl Jug Honorary Issue

M. Krack

E-mail: krack@phys.chem.ethz.ch

relativistic effects can be included into the pseudopotential parametrization. There are only a few properties like the nuclear magnetic resonance for which an all-electron treatment is inevitable. Also plane waves methods require to employ pseudopotentials, since an expansion of an atomic all-electron density or wavefunction in plane waves is computationally inefficient. Thus, there was always a strong need for efficient pseudopotentials from the very beginning.

This work refers to the relativistic, norm-conserving, separable, dual-space pseudopotentials of Goedecker, Teter, and Hutter (GTH) [1,2]. These are accurate pseudopotentials which have a compact analytic form based on Gaussian functions, whereas many other pseudopotential types are defined by dense radial grids which have to be tabulated for each atomic kind [3]. The GTH pseudopotential parameter sets for the elements from H to Rn optimized in the framework of the local density approximation (LDA) are already available from [2]. The original idea was to provide only optimized LDA parameter sets for all chemically relevant elements together with the code for the optimization. However, there is still no easy and foolproof recipe for the generation of new parameter sets and an optimization might be cumbersome, since there are often many pitfalls. While the LDA is still often used in condensed matter applications, nowadays most studies of, for instance, (bio-)molecular systems are performed with exchange-correlation functionals based on the generalized gradient approximation (GGA). The aim of this work is to provide the optimized GTH parameter sets for the elements from H to Kr for a couple of often used GGA functionals.

#### 2 Definition of the GTH pseudopotentials

The norm-conserving, separable, dual-space GTH pseudopotentials consist of a local part

$$V_{\text{loc}}^{\text{PP}}(r) = -\frac{Z_{\text{eff}}}{r} \operatorname{erf}\left(\alpha^{\text{PP}}r\right) + \sum_{i=1}^{4} C_{i}^{\text{PP}}\left(\sqrt{2}\alpha^{\text{PP}}r\right)^{2i-2} \exp\left[-\left(\alpha^{\text{PP}}r\right)^{2}\right], \quad (1)$$

Computational Science, Department of Chemistry and Applied Biosciences, ETH Zurich, USI-Campus, via G. Buffi 13, CH-6900 Lugano, Switzerland

with

$$\alpha^{\rm PP} = \frac{1}{\sqrt{2}r_{\rm loc}^{\rm PP}},$$

$$V_{\rm nl}^{\rm PP}(\mathbf{r},\mathbf{r}') = \sum_{lm} \sum_{ij} \langle \mathbf{r} \mid p_i^{lm} \rangle h_{ij}^l \langle p_j^{lm} \mid \mathbf{r}' \rangle, \qquad (2)$$

with the Gaussian-type projectors

$$\langle \mathbf{r} \mid p_i^{lm} \rangle = N_i^l Y^{lm}(\hat{r}) r^{l+2i-2} \exp\left[-\frac{1}{2} \left(\frac{r}{r_l}\right)^2\right],$$

where the  $N_i^l$  are normalization constants and the  $Y^{lm}(\hat{r})$  are spherical harmonics. An advantage of such a pseudopotential definition in real space is that it has an analytic representation in Fourier space, too. This feature is advantageous for methods dealing with both real and Fourier space like plane waves methods. The explicit form of Eqs. (1) and (2) in Fourier space is given in [2]. The attribute dual-space for the GTH pseudopotential stems from this special property. Moreover, a good convergence behavior is achieved both in real and Fourier space by the usage of Gaussian functions. A further advantage of the Gaussian functions is that with atom-centered Gaussian function basis sets the pseudopotential contribution involves only the calculation of two- and three-center overlap integrals over Gaussian functions. The three-center overlap integrals of Eq. (2) reduce to products of two-center overlap integrals because of the separability of the non-local term. There are very efficient recursion schemes [4] to evaluate such integrals. For that reason, the GTH pseudopotentials are optimally suited for the use with the Gaussian plane waves (GPW) method [5–7] which uses Gaussian basis sets for the expansion of the Kohn-Sham orbitals and an auxiliary plane waves basis set for the description of the electronic density.

Relativistic all-electron wavefunctions are used as the reference for the fit of the pseudo wavefunctions and hence all orbitals with an angular momentum quantum number l > 0split up into a spin-up and a spin-down orbital with the total angular momentum numbers  $j = l \pm 1/2$  and thus two different potentials are obtained. A weighted average potential

$$V_{l}(\mathbf{r}, \mathbf{r}') = \frac{1}{2l+1} \left[ l V_{l-\frac{1}{2}}(\mathbf{r}, \mathbf{r}') + (l+1) V_{l+\frac{1}{2}}(\mathbf{r}, \mathbf{r}') \right]$$
(3)

was defined by Bachelet and Schlüter [8]. Then the spin-orbit (SO) coupling is given by the difference potential

$$\Delta V_{l}^{\rm SO}(\mathbf{r}, \mathbf{r}') = \frac{2}{2l+1} \left[ V_{l+\frac{1}{2}}(\mathbf{r}, \mathbf{r}') - V_{l-\frac{1}{2}}(\mathbf{r}, \mathbf{r}') \right].$$
(4)

of the two-spin components. In this way the total pseudopotential can be rewritten as a sum

$$V(\mathbf{r}, \mathbf{r}') = V_{\text{loc}}(r) \,\delta(\mathbf{r} - \mathbf{r}') + \sum_{l} V_{l}(\mathbf{r}, \mathbf{r}') + \Delta V_{l}^{\text{SO}}(\mathbf{r}, \mathbf{r}') \,\mathbf{L} \cdot \mathbf{S}, \qquad (5)$$

of a scalar relativistic part and the difference potential  $\Delta V_l^{SO}$ (**r**, **r**'). The last term of Eq. (5) is neglected and an average scalar relativistic pseudopotential is obtained which has the same analytic form as the nonrelativistic pseudopotential given by Eqs. (1) and (2).

Table	1	Goedecker,	Teter	and	Hutter	pseudopotential	parameters
optimi	zeo	d for BLYP					-

Н	1	0.200000	-4.195961	0.730498		
He	2	0.200000	-9.147371	1.711978		
Li	3	0.400000	-14.116798	9.677248	-1.798865	0.086250
Be	4	0.325000	-24.089771	17.302757	-3.345610	0.166015
В	3	0.424072	-6.108908	0.985870		
		0.370860	6.343396			
С	4	0.338066	-9.136269	1.429260		
		0.302322	9.665512			
Ν	5	0.282871	-12.736467	1.951079		
		0.255234	13.678932			
0	6	0.243420	-16.991892	2.566142		
		0.220831	18.388851			
F	7	0.213840	-21.902415	3.275976		
-		0.194590	23,796583			
Ne	8	0.190000	-28.619598	4,155495		
1.0	0	0 178238	27 957849	0.833656		
		0.170200	21.957019	-1.076245		
		0 152764	0 331170	1.070245		
Na	9	0.233965	-2 689483	_0 509478		
144	/	0.233703	32 857159	0.507470		
		0.173100	_13 000008			
Ma	10	0.123199	-10.330203	2 860058		
wig	10	0.200984	40 674730	2.000030		
		0.141060	10 726171			
Δ1	2	0.104301	7 666832			
AI	5	0.450000	-7.000832	1 593970		
		0.497730	0.403032	-1.383870		
		0 565458	1 818267	2.044707		
C;	4	0.303438	6 250587			
51	4	0.440000	-0.239387	2 222770		
		0.444032	8.314009	-2.552779		
		0 502702	2 222419	5.011005		
р	5	0.302792	2.332410			
P	3	0.430000	-3.929333	2 121600		
		0.402809	10.414839	-5.151088		
		0 452459	2 052592	4.042991		
C	~	0.452458	2.953583			
5	6	0.420000	-5.988800	2 01 200 5		
		0.369/33	12.556485	-3.912905		
		0.412076	2 507721	5.051538		
<b>C1</b>	-	0.4128/6	3.587721			
CI	1	0.410000	-6.357052			
		0.343119	14.681348	-4.640343		
				5.990657		
		0.381135	4.222924			
Ar	8	0.400000	-7.100000			
		0.321550	16.800999	-5.328711		
				6.879336		
		0.355444	4.877196			
Κ	9	0.400000	-2.880134	-1.211435		
		0.306347	17.510023	-5.610379		
				7.242968		
		0.321058	6.903211	-2.199258		
				2.602197		

#### 3 Optimization of the GTH pseudopotential parameters

Firstly, the atomic all-electron wave functions for the actual electronic ground-state configurations were obtained from fully relativistic density functional calculations using a numerical atomic program. These all-electron wave functions were generated for each GGA functional and served as a reference for the optimization of the pseudo wavefunctions. The optimization of GTH pseudopotentials involves by definition

 Table 2 Goedecker, Teter and Hutter pseudopotential parameters optimized for BLYP

 Table 3 Goedecker, Teter and Hutter pseudopotential parameters optimized for BLYP

Ca	10	0.390000	-4.035611	-1.612574	Cu	11	0.530000			
		0.289365	20.425660	-7.236748			0.430782	10.298526	-6.058370	1.700546
		0.011550		9.342602					10.587260	-4.390790
		0.311779	7.531694	-2.085377			0 550005	2 744597	0.962055	3.485082
		0.601410	0.055223	2.407451			0.550805	2.744587	-0.862955	
Sc	11	0.376875	9.840097	-1 165860			0.265586	-12 661582	1.021002	
30	11	0.359989	3 306103	2 547964	Zn	12	0.510000	-12.001382		
		0.557707	5.500105	-3.289408	2.11	12	0.398550	11.959460	-8.665221	2.808077
		0.220994	-0.434882	6.464258					15.754090	-7.250424
				-7.648614						5.754846
		0.233380	-10.134039				0.545562	2.510500	-0.440325	
Ti	12	0.380000	8.711442	-0.700287					0.520999	
		0.337771	2.575264	3.692971	~		0.251685	-14.323098		
		0.040504		-4.767605	Ga	13	0.490000	10.0501.00		0 (00 (00
		0.242531	-4.630541	8.870875			0.414929	10.8/9160	-4.784493	0.608688
		0.242217	0 406652	-10.496161					/.166/85	-1.5/1626
v	12	0.245517	-9.400033	0 222250			0 567421	1 780264	0 277278	1.247439
v	15	0.373000	1 992040	-0.522259			0.307431	1.780204	-0.328198	
		0.527045	1.772040	-6.118233			0 238927	-16.078668	0.520170	
		0.246213	-5.549745	8.764202	Ge	4	0.540000	10.070000		
				-10.369943		-	0.427374	7.556515	-0.070902	-1.725963
		0.241153	-9.444362						-2.715745	4.456418
Cr	14	0.370000	5.814981	-0.675718						-3.537172
		0.316714	2.606983	5.067158			0.574899	0.800563	0.713074	
				-6.541672					-0.843721	
		0.239613	-4.666682	7.572902		-	0.785473	0.218708		
			11 10000	-8.960379	As	5	0.520000		0.445550	0.000.400
	1.5	0.220929	-11.189892	0 444510			0.456983	5.5/9/64	0.117773	-0.992439
Mn	15	0.365000	0.1/8/03	-0.444510					-1./66246	2.562466
		0.297644	1.094550	-8 250144			0 554947	1 000977	0.683251	-2.055895
		0 245825	-6 537424	7 845912			0.554947	1.000977	-0.808433	
		0.245025	0.557424	-9.283408			0.684981	0.348181	0.000455	
		0.222447	-11.669616	2.200.100	Se	6	0.510000	010 10101		
Fe	16	0.360000	7.011025	-0.222747			0.433675	6.433699	-0.220640	-1.175453
		0.276205	0.611496	7.992357					-1.627030	3.035007
				-10.318089						-2.408962
		0.247130	-8.710556	8.684169			0.472483	2.239703	0.401919	
				-10.275248				· · · · · · · -	-0.475556	
C	17	0.223534	-12.411504	0 (14012	р	7	0.609118	0.494947		
Co	17	0.355000	3.432978	0.614913	Br	1	0.500000	( 221282	0.219902	1 124(7)
		0.275554	0.193629	9.181280			0.436123	0.231283	0.218893	-1.1240/0
		0 243128	-8 922049	9 232630					-2.111875	2.905902
		0.243120	-0.922049	-10.924196			0 453793	2 465743	0 496368	-2.304901
		0.223625	-11.758783	10.924190			0.455775	2.405745	-0.587311	
Ni	18	0.350000	4.076029	0.233128			0.533156	0.747241		
		0.257147	0.098785	10.148642	Kr	8	0.500000			
				-13.101841			0.423621	6.373829	0.498586	-1.419917
		0.249753	-8.348615	7.266771					-2.953167	3.666210
				-8.598160						-2.909964
		0.212478	-13.672933				0.433913	2.632304	0.645177	
							0 512154	0 700545	-0./63384	
							0.313134	0.709343		

147

only a small set of parameters as it can be retrieved from Eqs. (1) and (2). The parameters are  $r_{loc}^{PP}$  and  $C_i^{PP}$  for the local part and  $r_l$  and  $h_{ij}^l$  for the non-local part. The GTH pseudopotential parameter sets for the elements from H to Rn optimized for LDA are already available from [2]. These parameter sets provided a good initial guess for the optimization of the GGA parameter sets and the same pattern of nonzero parameters was used for the fit of the corresponding GGA parameter sets, whereas the weights for the penalty functions were adapted in

order to achieve an optimal fit for all pseudo wavefunctions. A stubborn but robust simplex-downhill procedure was used for the optimization. The typical difference between the optimized pseudo wavefunction and corresponding all-electron reference wavefunction in the valence region was usually less than  $10^{-5}$  for the occupied valence orbitals and  $10^{-3}-10^{-4}$ 

 Table 4 Goedecker, Teter and Hutter pseudopotential parameters optimized for BP

Н	1	0.200000	-4.185769	0.726940			Ca	10	0.390000	-5.150665	-1.276359
He	2	0.200000	-9.130075	1.705475					0.291806	20.375018	-6.392254
Li	3	0.400000	-14.060569	9.585899	-1.772485	0.084552					8.252364
Be	4	0.325000	-24.051797	17.250016	-3.333976	0.165569			0.305851	8.086791	-2.351948
В	3	0.416891	-5.922294	0.924562							2.782862
		0.371077	6.295996						0.690846	0.050575	
С	4	0.336797	-8.868487	1.354476			Sc	11	0.385000	8.151138	-0.545788
		0.302410	9.622712						0.364086	2.633743	3.011895
Ν	5	0.283092	-12.450889	1.875196							-3.888340
		0.255290	13.636569						0.243373	-2.596433	7.977854
0	6	0.244018	-16.705908	2.492927							-9.439525
		0.220865	18.347594						0.253163	-8.161687	
F	7	0.214489	-21.618322	3.205432			Ti	12	0.380000	8.752405	-0.720513
		0.194610	23.757123						0.337256	2.577937	3.671930
Ne	8	0.190000	-27.056031	4.335635							-4.740441
		0.176097	28.150757	0.833656					0.242447	-4.635718	8.847008
				-1.076245							-10.467920
		0.193497	-0.240663						0.243277	-9.408158	
Na	1	0.754736	-2.139853				V	13	0.375000	7.151792	-0.343565
		0.722073	2.089328	-0.338912					0.329787	1.967060	4.726242
				0.437533							-6.101552
		0.875320	0.497225						0.246174	-5.582364	8.796910
Na	9	0.210332	1.692589	0.503580							-10.408644
		0.131731	40.129336						0.241159	-9.447430	
		0.140673	-12.691904				Cr	14	0.370000	5.658643	-0.680989
Mg	10	0.203445	-19.148633	2.823116					0.314507	2.825352	5.001950
		0.141013	40.615281								-6.457490
	•	0.104856	-10.812360						0.240791	-4.437221	7.252682
AI	3	0.450000	-7.517924	1.01.5.00						11 10 5 100	-8.581489
		0.486266	6.977756	-1.915682				1.5	0.220357	-11.185498	0 45560
		0.5(1000	1.0550.40	2.473135			Mn	15	0.365000	6.0869/3	-0.45/6/9
<u>.</u>		0.561928	1.855842						0.296649	1.826416	6.3/3014
51	4	0.440000	-6.31/841	2 (040(2					0.04(011	( 51555)	-8.22/525
		0.436036	8.904625	-2.694063					0.246911	-0.515555	7.910510
		0 409 410	2 410275	3.478020					0 222259	11 (10204	-9.359842
п	E	0.498419	2.4193/5				Б.	16	0.222558	-11.018284	0 220(05
P	3	0.430000	-3.892290	2 459022			ге	10	0.300000	0./002/1	-0.229003
		0.396816	10.932888	-3.458033					0.278019	0.640925	10 202870
		0 440121	2 020049	4.404501					0.252111	7.012426	-10.2028/0
c	6	0.449121	5.050048						0.232111	-7.913420	/.000133
3	0	0.420000	-0.055042	4 217067					0 222865	12 287740	-9.070093
		0.303400	15.051205	-4.217007			Co	17	0.222803	-12.367749	0 600787
		0.410245	3 678220	5.444211			CO	17	0.333000	0.608700	0.009787
C1	7	0.410243	6 304602						0.275447	0.008700	11 605801
CI	/	0.410000	-0.394002 15 113473	_4 008115					0 244865	-8 731042	0 10//18
		0.540018	15.115475	6 3 3 6 3 4 0					0.244005	-0.751942	10 979093
		0 3701/1	1 306336	0.550549					0 223214	-11 659403	-10.070905
٨r	8	0.377141	-7 100000				Ni	18	0.225214	1 080020	0.678735
Л	0	0.400000	17 158055	_5 553322			111	10	0.261517	0.640514	0.078755
		0.517220	17.150055	7 169309					0.201017	0.040514	-12 872890
		0 353911	4 943474	7.107507					0 222997	-11 035849	12 324159
K	9	0.400000	-3.208037	-1.139580					0.222771	11.055077	-14 582141
	-	0.305317	17.821132	-5.624592					0.215316	-12.607378	1
		5.000017	1	7.261316					0.210010	12.007070	
		0.317286	7,270738	-2.458363							
				2.908774							

for the unoccupied orbitals. Thus all optimizations were performed on a level corresponding to the LDA parameter sets.

In that way the parameters for the elements from H to Kr were optimized for the gradient-corrected exchange-correlation potentials of Becke, Lee, Yang, and Parr (BLYP) [9–11], Becke and Perdew (BP) [9,12], and Perdew, Burke and Ernzerhof (PBE) [13]. Only the semi-core versions of the

pseudopotentials which include partially the core electrons were optimized, since the experience with the LDA pseudopotentials showed that only these pseudopotentials are very accurate in most cases.

The optimized parameter sets obtained for the BLYP, BP, and PBE functional are listed in the Tables 1–9.

The table entries refer to the following format:

mize	d for B	BP	er une riener pot	europotentiul p	and the option	miz	ed fo	or PBE	, 10001 4110 110	uer poeudope	paran	inerens opti
Cu	11	0.530000				Н	1	0.200000	-4.178900	0.724463		
		0.428260	9.929973	-6.733781	2.061234	He	2	0.200000	-9.122144	1.702708		
				12.045870	-5.322083	Li	3	0.400000	-14.081155	9.626220	-1.783616	0.085152
					4.224272	Be	4	0.325000	-24.067467	17.279022	-3.339106	0.165549
		0.561800	2.527065	-0.760266 0.899559		В	3	0.418991 0.371320	-5.859462 6.297280	0.903756		
Zn	12	0.264818 0.510000	-12.790290	0.077007		С	4	0.338471	-8.803674	1.339211		
211	12	0.398535	11.714302	-8.952715	3.103896	Ν	5	0.283791	-12.415226	1.868096		
				16.606392	-8.014224 6.361093	0	6	0.255405 0.244554	-16.667215	2.487311		
		0.542780	2.594550	-0.564109 0.667463		F	7	$0.220956 \\ 0.214930$	$18.337458 \\ -21.573028$	3.199776		
Ga	13	0.251199 0.490000	-14.423696			Ne	8	0.194684 0.190000	23.743540 - 27.120160	4.360450		
		0.414448	10.636900	-5.036262	0.717210			0.176059	28.177371	0.833656		
				7.668329	-1.851829					-1.076245		
					1.469844			0.195475	-0.236294			
		0.570055	1.755905	0.262394		Na	9	0.236523	0.295105	-0.913885		
				-0.310469				0.143560	34.601492			
		0.238636	-16.154283					0.129932	-14.277462			
Ge	4	0.540000				Mg	10	0.192758	-20.575391	3.040167		
		0.426527	7.460088	-0.101485	-1.799778			0.141407	41.047292			
				-2.796322	4.647007			0.102932	-9.985626			
					-3.688447	Mg	2	0.576960	-2.690407			
		0.570818	0.832322	0.672048 - 0.795178				0.593924	3.503211	-0.716772 0.925348		
		0.802293	0.201406					0.707157	0.831158			
As	5	0.520000				Al	3	0.450000	-7.554761			
		0.456157	5.518370	0.173127 -2.081309	-1.142209 2.949170			0.487435	6.959938	-1.888836 2.438477		
					-2.340831			0.562189	1.865299			
		0.556427	0.968716	0.692518		Si	4	0.440000	-6.269288			
				-0.819399				0.435634	8.951742	-2.706271		
c	~	0.700852	0.313103					0 407040	0 101077	3.493781		
Se	6	0.510000	( 107(00	0.11(((0)	1 250025	D	~	0.49/942	2.4312//			
		0.433/81	6.42/602	-0.116660	-1.250925	Р	5	0.430000	-5.8/5943	2 470256		
				-1.884023	-2.563634			0.390377	11.008802	4.480210		
		0.472877	2.193509	0.463547				0.448298	3.056064			
				-0.548477		S	6	0.420000	-5.986260			
		0.617793	0.457876					0.364820	13.143544	-4.241830		
Br	7	0.500000								5.476180		
		0.438443	6.028091	0.369508	-1.265836		_	0.409480	3.700891			
				-2.535916	3.268375	Cl	7	0.410000	-6.392082	4 00 4500		
		0 452(52	2 45 4 4 40	0.552264	-2.594192			0.339539	15.218990	-4.934523		
		0.452655	2.454449	0.555204				0 278 474	1 220775	0.370442		
		0 572701	0 534532	-0.034031		٨r	8	0.378474	-7 100000			
Kr	8	0.500000	0.554552			лі	0	0.400000	17 252038	-5 585488		
	0	0.422012	6.436401	0.554844	-1.490192			0.010010	17.252050	7.210834		
		0.122012	01120101	-3.140099	3.847659			0.353370	4.974216			
					-3.053985	Κ	9	0.400000	-3.363552	-1.086530		
		0.433814	2.587592	0.721291				0.305318	17.850623	-5.622649		
				-0.853443						7.258808		
		0.547605	0.528788					0.316484	7.333780	-2.460945		
										2.911829		
Eler	nent	Zeff	$r_{100}$ $C_1$	$C_2$	$C_2 = C_4$			rə	$h_{1}^{2}$	$h_{1,2}^2$	$h_{1}^{2}$	1
		-en	$h_{\rm r}$	$k^0$	$L_{0}^{0}$			2	1,1	$h^{\frac{1}{2},2}$	$h^{1,3}$	,
			$r_0 n_{1,1}$	$n_{1,2}$	<i>i</i> <sub>1,3</sub>					<i>n</i> <sub>2,2</sub>	<sup>11</sup> 2,3	
				$h_{2,2}^0$	$h_{2,3}^0$						$h_{3,3}^2$	1
				,_	$h_{3,3}^{0}$	Or	1.7 +1	na naramat	ore which w	ere ontimiz	ad are lista	d for each
			$r_1$ $h_1^1$	$h_{1}^1$	$h_{1,2}^{\tilde{1},\tilde{2}}$		1 y U mar		$x_{10} = x_{10} = x$	ore optimized	the off diag	a tot caell
			· 1 "1,1	$h^{1,2}$	-1,3 1	ele	mer	n. martwig			ne on-diag	
				<sup>11</sup> 2,2	<sup>2</sup> 2,3	me	nts	of the sym	metric coeff	cient matri	$\mathbf{x} \mathbf{n}$ for the	non-local

 $h^0_{1,3} \\ h^0_{2,3} \\ h^0_{3,3} \\ h^1_{1,3} \\ h^1_{2,3} \\ h^1_{3,3}$ 

 Table 6
 Goedecker, Teter and Hutter pseudopotential parameters optimized for BP

Only the parameters which were optimized are listed for each element. Hartwigsen et al. [2] derived the off-diagonal elements of the symmetric coefficient matrix  $\mathbf{h}^l$  for the non-local projectors from the diagonal elements using a set of explicit

149

Table 7 Goedecker, Teter and Hutter pseudopotential parameters opti-

Ca	10	0.390000	-4.167072	-1.583798	Cu	11	0.530000			
		0.289356	20.531876	-7.129786			0.431355	9.693805	-6.470165	1.935952
				9.204514					11.501774	-4.998607
		0.327882	5.805605	-0.428753						3.967521
		0 (70(17	0.050060	0.507308			0.561392	2.545473	-0.784636	
C	11	0.6/961/	0.058068	0 557050			0.064555	10.000(14	0.928394	
Sc	11	0.385000	8.214900	-0.55/059	7.	10	0.264555	-12.828614		
		0.303011	2.040555	3.021084	Zn	12	0.510000	11 520041	9 701909	2 145096
		0 243909	2 624822	-3.900203			0.400310	11.550041	-6.791898	5.145080 9.120579
		0.243696	-2.034623	0.456424					10.403773	-0.120370
		0 253206	-8 1650/8	-9.450424			0 5/3182	2 507105	-0 594263	0.445509
Ті	12	0.380000	8 711442	-0.700287			0.545162	2.397193	0 703141	
11	12	0.337771	2 575264	3 692971			0.250959	-14 466958	0.705141	
		0.557771	2.575204	-4767605	Ga	13	0.490000	14.400/50		
		0 242531	-4 630541	8 870875	Ou	10	0.413962	10 592918	-5 152803	0 775294
		0.2 12331	1.050511	-10496161			0.113702	10.372710	7 913255	-2.001799
		0.243317	-9.406653	101190101					1010200	1.588879
V	13	0.375000	7.474704	-0.370264			0.570154	1.759998	0.251224	
		0.327795	1.940878	4.725688					-0.297253	
				-6.100837			0.238368	-16.211820		
		0.244766	-5.978167	9.358639	Ge	4	0.540000			
				-11.073291			0.421865	7.510241	-0.588108	-1.447976
		0.241739	-9.499891						-1.595888	3.738657
Cr	14	0.370000	5.699658	-0.695486						-2.967467
		0.313934	2.869955	4.974456			0.567529	0.913860	0.546875	
				-6.421996					-0.647072	
		0.240866	-4.476209	7.331414			0.813914	0.197177		
				-8.674646	As	5	0.520000			
		0.220286	-11.197116				0.455550	5.520673	0.035122	-1.061082
Mn	15	0.365000	6.093046	-0.446469					-1.771193	2.739703
		0.295686	1.88/120	6.356837			0.554606	1 001700	0 (20200	-2.174572
		0.045(12	6 570025	-8.206641			0.554606	1.021792	0.629208	
		0.245613	-6.5/0025	7.983360			0.702(90	0.214705	-0./44489	
		0 222522	11 612051	-9.446039	<b>C</b> •	6	0.703089	0.314795		
Fo	16	0.222323	-11.012031	0 228833	36	0	0.310000	6 5 1 8 1 1 0	0 222716	1 106120
ге	10	0.300000	0.730789	-0.220033			0.432400	0.518110	-0.222710	-1.190129
		0.278203	0.029300	-10215810					-1.03/9/8	-2 451335
		0 251383	_7 032133	7 607070			0 470402	2 281262	0 365335	-2.451555
		0.251505	-7.952155	-9107307			0.470492	2.201202	-0.432271	
		0 222856	-12 385799	2.107507			0.625600	0 439799	0.452271	
Ni	18	0.350000	2.102166	0.648484	Br	7	0.500000	0.157777		
	10	0.261295	0.622658	9.970227	21		0.438039	6.078556	0.330498	-1.238382
				-12.871507					-2.440900	3,197489
		0.224253	-11.142708	12.429566						-2.537928
				-14.706861			0.453136	2.459304	0.522753	
		0.215348	-12.628146						-0.618530	
							0.567714	0.559266		
equati	ions and	thus they listed	l only the diagona	l elements <i>h</i> <sup>l</sup>	Kr	8	0.500000			
equali			i only the diagona				0.421657	6.465304	0.538660	-1.502601

 Table 8 Goedecker, Teter and Hutter pseudopotential parameters optimized for PBE

 Table 9
 Goedecker, Teter and Hutter pseudopotential parameters optimized for PBE

equations and thus they listed only the diagonal elements  $h_{ii}^{t}$ . In this work the coefficient matrix was orthonormalized using the matrix transformation

$$\mathbf{h}_{d}^{l} = \mathbf{U} \, \mathbf{h}^{l} \, \mathbf{U}^{\mathrm{T}} \quad \text{with} \quad \mathbf{U} \, \mathbf{U}^{\mathrm{T}} = \mathbf{1} \,, \tag{6}$$

where  $\mathbf{h}_{d}^{l}$  is a diagonal matrix. In that way only the contributions of the diagonal elements  $h_{ii}^{l}$  have to be computed during the optimization. On the other hand, now the upper triangle including the diagonal elements of the coefficient matrix  $\mathbf{h}^{l}$  has to be listed. The corresponding coefficient matrix for the SO term is not listed, but the parameters can be retrieved from the online database of the CP2K project [14], if an explicit calculation of the SO coupling is required and available.

#### 4 Test calculations

0.433744

0.524691

The optimized GTH pseudopotentials were tested using a set of small molecules. The plane waves code CPMD (version 3.9.1) [15] was used to optimize the structure of these small molecules. To this end, each molecule was placed in a simple

2.601165

0.635595

-3.139389

 $0.705110 \\ -0.834297$ 

3.879700

-3.079416



Fig. 1 Comparison of the bond distances calculated with CPMD [15] using the GTH pseudopotentials optimized for BLYP and the bond distances obtained by all-electron calculations with the 6-311++G(3df,3pd) basis set using the Gaussian 03 program [18] for the compounds of Tables 10 and 11 containing only the elements from H to Ar

cubic box of edge length 12 Å using the molecule option of CPMD, which allows to deal with isolated molecules, since the interaction with the periodic images becomes negligible. The convergence criterion for the atomic forces was set to 0.0001 a.u. for LDA and to 0.0005 a.u. for the GGA functionals. A wavefunction cutoff of 500 Ry for LDA and 1000 Ry for the GGA functionals was employed. Table 10 and 11 compare the structure data obtained with the new GTH pseudopotential parameter sets to the corresponding experimental data. The results for the GGA functionals show in most cases the typical bond elongation compared to the LDA data, whereas differences among the GGA functionals are rather small. The same trends are well-known from all-electron calculations. Molecules containing the elements from Sc to Ga were not included into the test set, since the requested convergence for the atomic forces could not be achieved because of the hardness of the semi-core pseudopotentials employed for these elements. However, these GTH pseudopotentials can be used with the Gaussian plane waves method as it is implemented in the QUICKSTEP code [5-7], which is part of the open source project CP2K [14]. QUICKSTEP is less affected by the hardness of these pseudopotentials, since the short-ranged integral terms of Eqs. (1) and (2) can be easily computed analytically over Gaussian functions (see Sect. 2). The development of Gaussian basis sets optimized for the use with the GTH pseudopotentials is in progress.

As a final test, the structures of the compounds of the Tables 10 and 11 were optimized at the all-electron level. The optimizations were performed with the Gaussian 03 program [18] employing the large standard Gaussian basis set 6-311++G(3df,3pd), which is only available for the elements from H to Ar. Thus just the compounds of the Tables 10 and 11 containing only the elements from H to Ar were consid-

 Table 10 Structure data of small molecules optimized using the GTH pseudopotentials

Molecule	[Å,°]	LDA	BLYP	BP	PBE	Exp.	Ref.
$H_2$	r(HH)	0.766	0.746	0.750	0.751	0.741	[16]
LiH	r(LiH)	1.603	1.599	1.603	1.603	1.596	[16]
LiF	r(LiF)	1.550	1.555	1.579	1.575	1.564	[16]
LiCl	r(LiCl)	2.000	2.031	2.033	2.023	2.021	[16]
LiBr	r(LiBr)	2.150	2.183	2.183	2.177	2.170	[16]
BeF <sub>2</sub>	r(BeF)	1.372	1.377	1.388	1.386	1.373	[17]
$BH_3$	r(BH)	1.198	1.190	1.196	1.196	1.190	[17]
BF <sub>3</sub>	r(BF)	1.310	1.333	1.328	1.327	1.313	[17]
BCl <sub>3</sub>	r(BCl)	1.727	1.755	1.748	1.746	1.742	[17]
BBr <sub>3</sub>	r(BBr)	1.883	1.920	1.911	1.907	1.893	[17]
$CH_4$	r(CH)	1.096	1.093	1.095	1.095	1.087	[17]
$C_2H_2$	r(CC)	1.226	1.226	1.226	1.226	1.203	[17]
	r(CH)	1.087	1.087	1.087	1.087	1.063	
$C_2H_4$	r(CC)	1.321	1.332	1.333	1.332	1.330	[17]
	r(CH)	1.093	1.087	1.091	1.091	1.080	
	a(HCH)	116.7	116.5	116.6	116.6	117.1	
$C_2H_6$	r(CC)	1.508	1.538	1.532	1.527	1.522	[17]
	r(CH)	1.100	1.096	1.099	1.099	1.089	
	a(CCH)	111.7	111.4	111.4	111.5	111.2	
CH <sub>3</sub> F	r(CH)	1.101	1.094	1.098	1.099	1.100	[17]
	r(CF)	1.376	1.418	1.403	1.401	1.383	
	a(HCH)	109.8	110.6	110.2	110.2	110.6	
CH <sub>3</sub> Cl	r(CH)	1.096	1.089	1.093	1.094	1.087	[17]
	r(CCl)	1.760	1.816	1.794	1.787	1.776	
	a(HCH)	110.0	110.8	110.5	110.4	110.4	
CH3OH	r(CH)	1.106	1.099	1.103	1.103	1.098	[17]
	r(CO)	1.406	1.446	1.433	1.430	1.429	
	r(OH)	0.971	0.971	0.970	0.969	0.975	
CO	r(CO)	1.125	1.134	1.135	1.135	1.128	[16]
$CO_2$	r(CO)	1.161	1.170	1.171	1.171	1.160	[17]
$N_2$	r(NN)	1.093	1.099	1.101	1.102	1.098	[16]
$N_2O$	r(NN)	1.129	1.136	1.137	1.137	1.127	[17]
	r(NO)	1.178	1.203	1.194	1.191	1.185	
$NH_3$	r(NH)	1.022	1.020	1.022	1.022	1.014	[17]
	a(HNH)	107.6	107.2	106.5	106.4	107.2	
HCN	r(CH)	1.078	1.071	1.073	1.074	1.065	[17]
	r(CN)	1.148	1.155	1.156	1.157	1.153	
$H_2O$	r(OH)	0.971	0.972	0.970	0.970	0.958	[17]
	a(HOH)	105.1	104.7	104.4	104.4	104.5	
$H_2O_2$	r(OO)	1.434	1.494	1.472	1.468	1.456	[17]
	r(OH)	0.978	0.978	0.977	0.976	0.967	
	a(OOH)	100.8	99.7	99.7	99.8	102.3	
_	d(HOOH)	112.8	111.0	110.5	113.7	113.7	
H2	r(FF)	1.388	1.434	1.417	1.415	1.412	[16]
HF	r(HF)	0.933	0.933	0.932	0.932	0.917	[16]
$OF_2$	r(OF)	1.394	1.442	1.425	1.422	1.405	[17]
	a(FOF)	104.2	104.3	104.4	104.4	103.1	

ered. A tight convergence criterion was applied for both the wavefunction and the structure optimizations and the use of any molecular symmetry was disabled. The exchange-correlation functional BLYP was employed using ultra-fine integration grids. The comparison of the all-electron and the GTH pseudopotential results is shown in Fig. 1.

The agreement between the all-electron and the GTH pseudopotential results is very satisfactory considering that Gaussian 03 and CPMD are based on quite different methods. Moreover, differences in the functional implementations may cause small deviations. The molecules LiF and NaH show the largest deviation with 0.029 Å, but most of the 56 bond distances differ only by less than 0.01 Å.

 Table 11
 Structure data of small molecules optimized using the GTH pseudopotentials

Molecule	[Å,°]	LDA	BLYP	BP	PBE	Exp.	Ref.
NaH	r(NaH)	1.866	1.855	1.901	1.903	1.887	[16]
NaF	r(NaF)	1.910	1.938	1.947	1.958	1.926	[16]
NaCl	r(NaCl)	2.332	2.378	2.379	2.382	2.361	[16]
NaBr	r(NaBr)	2.468	2.496	2.522	2.523	2.502	[16]
AlH	r(AlH)	1.670	1.660	1.669	1.677	1.648	[16]
AlF <sub>3</sub>	r(AlF)	1.606	1.632	1.630	1.630	1.630	[17]
AlCl <sub>3</sub>	r(AlCl)	2.050	2.081	2.075	2.074	2.068	[17]
SiH <sub>4</sub>	r(SiH)	1.486	1.480	1.486	1.489	1.471	[17]
SiO	r(SiO)	1.497	1.510	1.515	1.517	1.510	[16]
PH <sub>3</sub>	r(PH)	1.427	1.424	1.427	1.429	1.413	[17]
-	a(HPH)	91.8	93.4	92.7	92.6	93.5	
HCP	r(CH)	1.082	1.074	1.079	1.080	1.066	[17]
	r(CP)	1.528	1.540	1.541	1.542	1.540	
PN	r(PN)	1.476	1.490	1.490	1.491	1.491	[16]
H <sub>2</sub> S	r(SH)	1.351	1.348	1.349	1.350	1.336	[17]
-	a(HSH)	91.2	92.1	91.7	91.6	87.8	
COS	r(CO)	1.159	1.169	1.169	1.169	1.154	[17]
	r(CS)	1.550	1.567	1.564	1.564	1.563	
CS	r(CS)	1.526	1.540	1.541	1.542	1.535	[16]
CS <sub>2</sub>	r(CS)	1.545	1.560	1.558	1.558	1.553	[17]
$SO_2$	r(SO)	1.428	1.447	1.443	1.444	1.431	[17]
2	a(OSO)	119.6	119.5	119.7	119.6	119.3	
$SF_6$	r(SF)	1.563	1.600	1.589	1.588	1.561	[17]
HCI	r(HCl)	1.289	1.289	1.288	1.288	1.275	[16]
KF	r(KF)	2.104	2.180	2.154	2.159	2.171	[16]
KCl	r(KCl)	2.598	2.684	2.665	2.659	2.667	[16]
KBr	r(KBr)	2.750	2.840	2.821	2.821	2.821	[16]
CaO	r(CaO)	1.781	1.822	1.810	1.812	1.822	[16]
CaS	r(CaS)	2.271	2.324	2.301	2.302	2.318	[16]
GeO	r(GeO)	1.592	1.623	1.620	1.631	1.625	[16]
GeH <sub>4</sub>	r(GeH)	1.514	1.520	1.525	1.534	1.514	[17]
As <sub>4</sub>	r(AsAs)	2.409	2.468	2.450	2.453	2.435	[17]
AsH <sub>3</sub>	r(AsH)	1.521	1.528	1.529	1.535	1.528	[17]
	a(HAsH)	90.6	91.9	91.3	91.0	91.9	
H <sub>2</sub> Se	r(SeH)	1.473	1.477	1.477	1.480	1.459	[17]
	a(HSeH)	90.0	90.7	90.4	90.3	91.0	
COSe	r(CO)	1.157	1.167	1.166	1.166	1.153	[17]
	r(CSe)	1.693	1.723	1.715	1.717	1.710	
CSe <sub>2</sub>	r(CSe)	1.682	1.706	1.702	1.704	1.692	[17]
SeO <sub>2</sub>	r(SeO)	1.608	1.635	1.630	1.633	1.608	[17]
	a(OSeO)	114.4	114.6	114.4	114.4	113.8	
HBr	r(HBr)	1.429	1.432	1.431	1.432	1.414	[16]
BrF	r(BrF)	1.758	1.771	1.789	1.788	1.759	[16]
BrCl	r(BrCl)	2.125	2.186	2.160	2.154	2.136	[16]
Br <sub>2</sub>	r(BrBr)	2.274	2.338	2.311	2.306	2.281	[16]

#### **5** Summary

Effective Gaussian-type pseudopotential parameter sets for the elements from H to Kr were presented. Parameter sets for the gradient-corrected exchange-correlation functionals BLYP, BP and PBE were optimized. Test calculation for small molecules were performed to show the accuracy and reliability of the presented pseudopotentials. All the GTH pseudopotential parameter sets for the presented exchange-correlation functionals including supplementary material are also available online [14]. The parameter sets are provided in different formats for a direct usage with the program packages CPMD and CP2K (QUICKSTEP).

Acknowledgements This work was partially supported by the Bundesministerium für Bildung and Forschung (BMBF, grant number 03N6015). The computer resources were provided by the Forschungszentrum Jülich (JUMP) and the Swiss National Supercomputing Center (CSCS) in Manno.

#### References

- 1. Goedecker S, Teter M, Hutter J (1996) Phys Rev B 54:1703
- 2. Hartwigsen C, Goedecker S, Hutter J (1998) Phys Rev B 58:3641
- 3. Troullier N, Martins JL (1991) Phys Rev B 43:1993
- 4. Obara S, Saika A (1986) J Chem Phys 84:3963
- 5. Lippert G, Hutter J, Parrinello M (1997) Mol Phys 92:477
- Krack M, Parrinello M In: High performance computing in chemistry, Report of the BMBF project, Grant Number 01IRA17 A-C, edited by Grotendorst J (Jülich FZ, Germany, 2004), vol. 25 of NIC series
- VandeVondele J, Krack M, Mohamed F, Parrinello M, Chassaing T, Hutter J (2005) Comput Phys Commun 167:103
- 8. Bachelet GB, Schlüter M (1982) Phys Rev B 25:2103
- 9. Becke AD (1988) Phys Rev A 38:3098
- 10. Lee CT, Yang WT, Parr RG (1988) Phys Rev B 37:785
- 11. Miehlich B, Savin A, Stoll H, Preuss H (1989) Chem Phys Lett 157:200
- 12. Perdew JP (1986) Phys Rev B 33:8822
- 13. Perdew JP, Burke K, Ernzerhof M (1996) Phys Rev Lett 77:3865
- 14. The CP2K developers group (2002) http://cp2k.berlios.de
- CPMD, Version 3.9.1, copyright IBM Corp. 1990–2004, copyright MPI für Festkörperforschung Stuttgart 1997–2001; http://www.cpmd.org
- 16. Huber KP, Herzberg G Constants of diatomic molecules, (data prepared by Gallagher JW, Johnson RD III) in NIST chemistry webBook, NIST standard reference database number 69, edited by Linstrom PJ, Mallard WG, March 2003, National institute of standards and technology, Gaithersburg MD, 20899 (http://webbook.nist.gov)
- 17. Graner G, Hirota E, Iijima T, Kuchitsu K, Ramsay DA, Vogt J, Vogt N Structure data of free polyatomic molecules. In: Kuchitsu K (ed) Landolt-Börnstein, New series, Group II: molecules and radicals, vol 25. Springer, Berlin Heidelberg New York
- Frisch MJ, Trucks GW, Schlegel HB, Scuseria GE, Robb MA, Cheeseman JR, Montgomery JA, Jr, Vreven T, Kudin KN, et al, (2003) Gaussian 03, Revision B.03, Gaussian, Inc. Pittsburgh