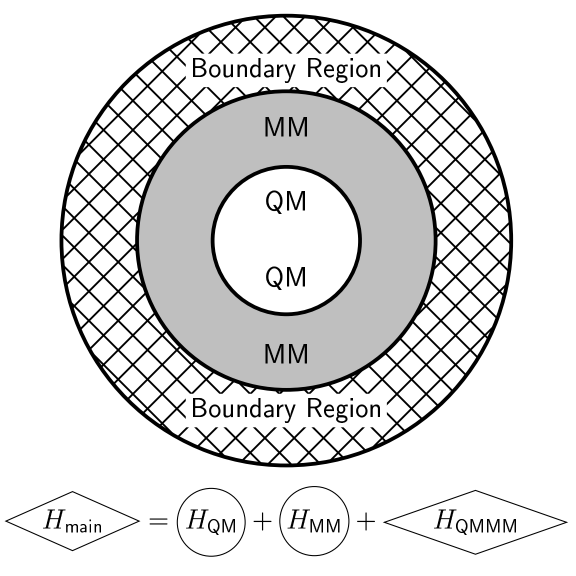
# Theory of hybrid Quantum Mechanical Molecular Mechanical methods An introduction to a way of treating complex chemical systems



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#### Outline of the talk

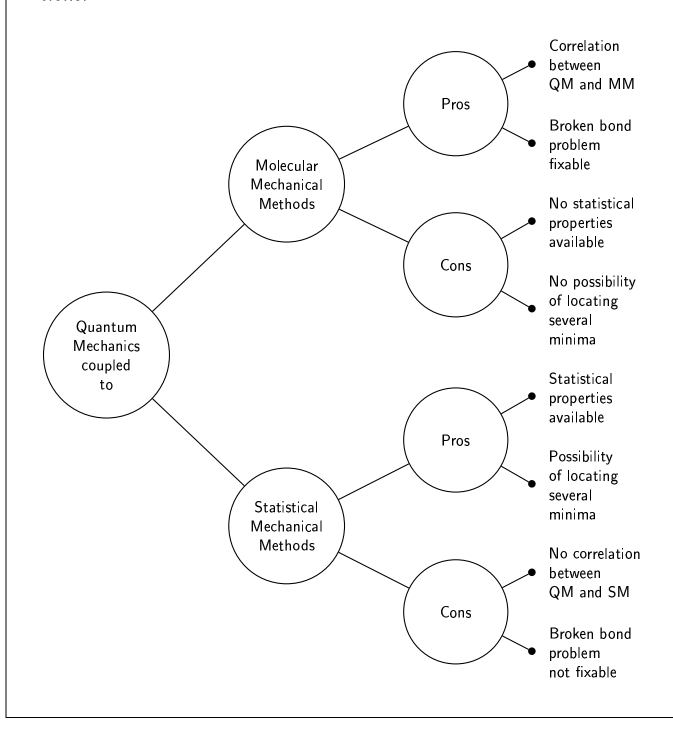
- Why couple QM and MM
- $\bullet$  Advantages and disadvantages of QM/MM in comparison to QM/SM
- Aspects of the hybrid QM MM method
- Criteria to fulfill when partitioning a molecular system into a QM part and an MM part
- How to treat broken bonds, if such are present after the partitioning
- Basics
- Establishing the main hamiltonian
- Example
- Acknowledgements
- Bibliography

### Why couple QM and MM

- For big molecules QM is to expensive in itself.
- No harm is done in using lower level theory far away from the chemically interesting place.
- If used with care, MM has proven its worth.
- Use QM coupling to MM as an alternative to QM coupling to SM.

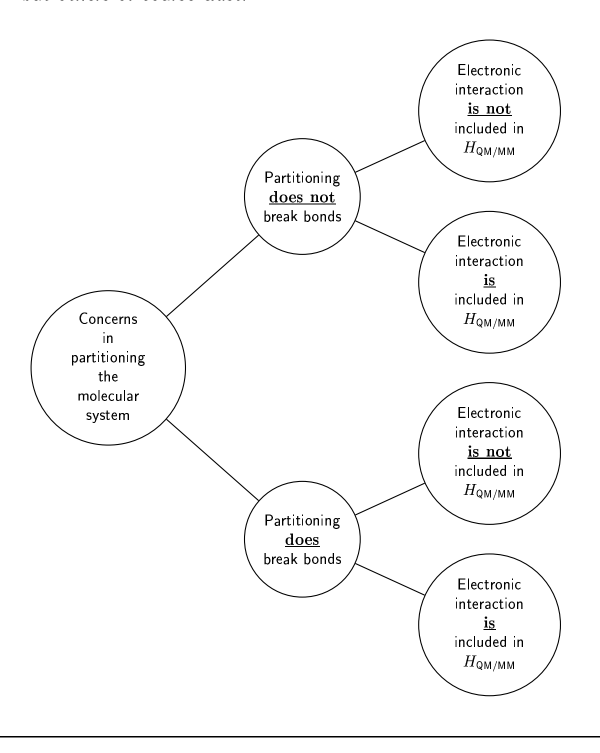
## Advantages and disadvantages of QM/MM in comparison to QM/SM

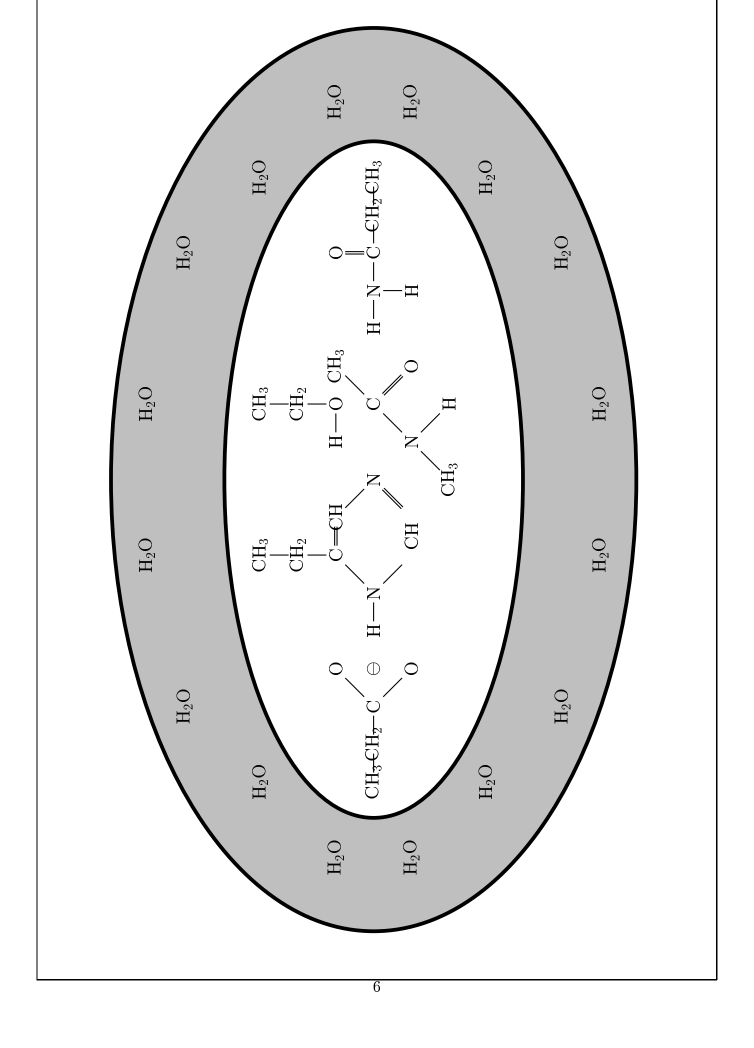
We list some <u>basic</u> pros and cons in comparing QM/MM and QM/SM, but stress, that both methods may be refined to annihilate the limitations.

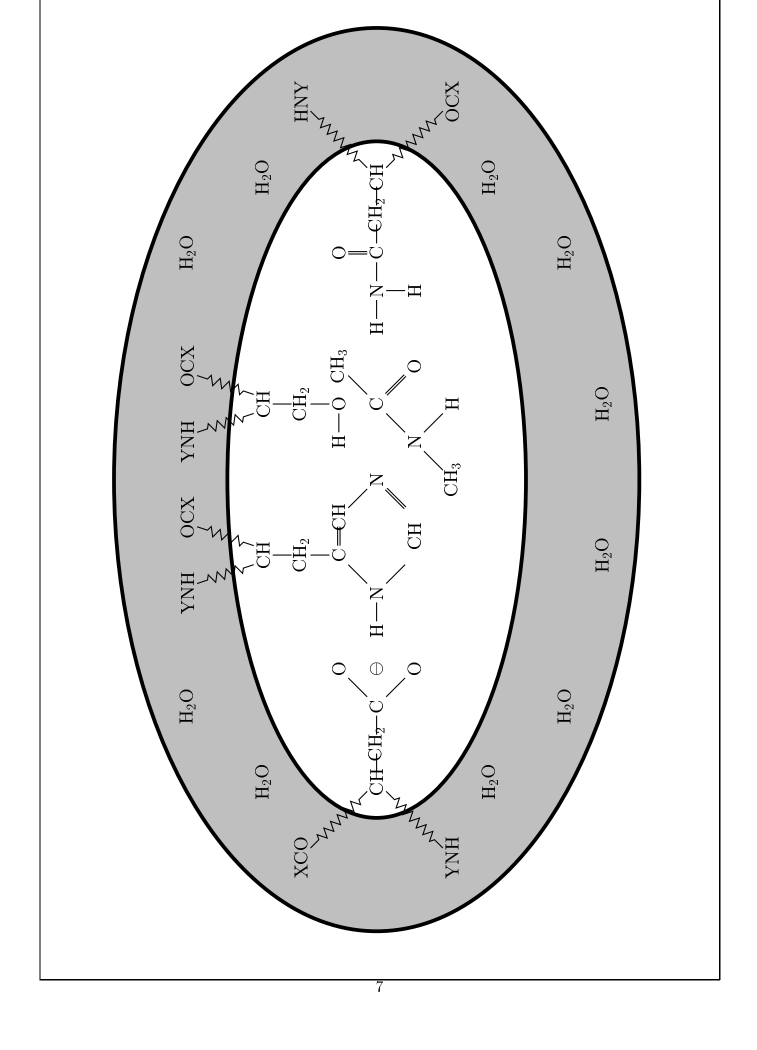


## Aspects of the hybrid QM MM method

We note the most <u>essential</u> aspects in the combined QM MM method, but others of course exist.







## Criteria to fulfill when partitioning a molecular system into a QM part and an MM part

- Fundamental knowledge of the system (Structure info in the case of enzymes).
- Largest possible size of QM part as opposed to level of accuracy necessary to treat QM part satisfactory.
- In case of calculating molecular properties, how will a broken bond between the QM region and the MM region affect the result.
- In case of modelling chemical reactions, how will the reaction affect the electronic distribution at the boundary (The goal is, that there should be no change in the electronic distribution).

## How to treat broken bonds, if such are present after the partitioning

- For each broken bond, a link atom should be added along the broken bond in order to be able to perform a QM calculation. The link atom is often chosen as a hydrogen atom and should not be allowed to interact with any MM atom.
- In addition to *nonbonded interaction terms* there are now also bonded interaction terms present.
- Bonded interaction terms: For each bond or angle or dihedral crossing the boundary, a force field stretch term or bend term or torsional term is added.
- Nonbonded interaction terms: Comparing [1] and [2] there seems to be some difference in approach.
  - [1]: Nonbonded interaction terms (van der walls interaction plus electrostatic interaction) between a QM atom and an MM atom are added if these are separated by more than three bonds!!!
  - [2]: Nonbonded interaction terms (van der walls interaction plus electrostatic interaction) between a QM atom and an MM atom are added whether or not these are separated by more than three bonds!!!

#### **Basics**

#### Choosing how to interact between QM and MM

- Comparing [1] and [2] there seems to be some difference in approach.

  - [2]: Method seems to be mainly based on treating  $\frac{\text{MM-partial-charge}}{\text{Consequence: Some terms from } H_{\text{QMMM}} \text{ will enter the HF SCF equations)}.$

#### Notation

- Indices *i*, *j* corresponds to QM <u>electrons</u>.
- Indices k, l corresponds to QM <u>nuclei</u>.
- Indices s, t corresponds to QM <u>atoms</u>.
- Indices m, n corresponds to MM <u>atoms</u>.
- Indices  $\alpha, \beta$  corresponds to basis functions.

Partitioning of the hamiltonian

$$H_{\mathsf{main}}\Psi = E\Psi$$

$$H_{\rm main} = H_{\rm QM} + H_{\rm MM} + H_{\rm QMMM}$$

$$E = \frac{\langle \Psi | H_{\text{main}} | \Psi \rangle}{\langle \Psi | \Psi \rangle}$$

$$E = E_{\rm QM} + E_{\rm MM} + E_{\rm QMMM}$$

Treating the separated parts

$$egin{array}{ll} H_{ extsf{QM}} & = & -rac{1}{2}\sum_{i}
abla_{i}^{2} - \sum_{i,k}rac{Z_{k}}{r_{ik}} + \sum_{i,j}rac{1}{r_{ij}} \ & + \sum_{k,l}rac{Z_{k}Z_{l}}{r_{kl}} \end{array}$$

$$\begin{split} H_{\text{MM}} &= \sum_{\text{bond}}^{\text{MM MM}} \frac{k_{\text{b}}}{2} (r - r_0)^2 + \sum_{\text{angle}}^{\text{MM MM}} \frac{k_{\text{a}}}{2} (\theta - \theta_0)^2 + \sum_{\text{dihedral}}^{\text{MM MM}} \frac{k_{\text{d}}}{2} (1 + \cos \Theta) \\ &+ \sum_{m,n} \left( \frac{A_{mn}}{r_{mn}^{12}} - \frac{B_{mn}}{r_{mn}^6} \right) + \sum_{\text{Hydrogen bond}} \left( \frac{C_{mn}}{r_{mn}^{12}} - \frac{D_{mn}}{r_{mn}^{10}} \right) \\ &+ \sum_{m,n} \frac{q_m q_n}{r_{mn}} \\ &+ \frac{1}{2} \sum_{m < n} q_m \frac{\boldsymbol{\mu}_n \cdot \boldsymbol{r}_{mn}}{r_{mn}^3} + \frac{1}{2} \sum_{m > n} q_m \frac{\boldsymbol{\mu}_n \cdot \boldsymbol{r}_{mn}}{r_{mn}^3} - \frac{1}{2} \sum_m \alpha_m \boldsymbol{E}_m \cdot \boldsymbol{E}_m \end{split}$$

$$H_{\mathsf{QMMM}}$$
 - Approach I - Based on [1] and [3]

One might at first try to not explicitly include electronic interactions in  $H_{\rm QMMM}$  thus getting

$$\begin{split} H_{\text{QMMM}} &= \sum_{\text{bond}}^{\text{QM MM}} \frac{k_{\text{b}}}{2} (r - r_0)^2 + \sum_{\text{angle}}^{\text{QM MM}} \frac{k_{\text{a}}}{2} (\theta - \theta_0)^2 + \sum_{\text{dihedral}}^{\text{QM MM}} \frac{k_{\text{d}}}{2} (1 + \cos \Theta) \\ &+ \sum_{s,m} \left( \frac{A_{sm}}{r_{sm}^{12}} - \frac{B_{sm}}{r_{sm}^6} \right) + \sum_{\text{Hydrogen bond}} \left( \frac{C_{sm}}{r_{sm}^{12}} - \frac{D_{sm}}{r_{sm}^{10}} \right) \\ &+ \sum_{s,m} \frac{q_s q_m}{r_{sm}} \\ &+ \frac{1}{2} \sum_{s,m} q_s \frac{\boldsymbol{\mu}_m \cdot \boldsymbol{r}_{sm}}{r_{sm}^3} \end{split}$$

This approach will make  $H_{\rm QMMM}$  independent on electronic coordinates, and thus  $H_{\rm QMMM}$  will just be a number to be added to the total energy.

One should in optimizations of the QM part add gradients from the MM part, as optimization of the QM part otherwise will proceed without correlation to the MM part.

$$H_{\mathrm{QMMM}}$$
 - Approach II - Based on [2] and [3]

One might then explicitly include electronic interactions in  $H_{\rm QMMM}$  thus getting

$$\begin{split} H_{\text{QMMM}} &= \sum_{\text{bond}}^{\text{QM MM}} \frac{k_{\text{b}}}{2} (r - r_0)^2 + \sum_{\text{angle}}^{\text{QM MM}} \frac{k_{\text{a}}}{2} (\theta - \theta_0)^2 + \sum_{\text{dihedral}}^{\text{QM MM}} \frac{k_{\text{d}}}{2} (1 + \cos \Theta) \\ &+ \sum_{s,m} \left( \frac{A_{sm}}{r_{sm}^{12}} - \frac{B_{sm}}{r_{sm}^6} \right) + \sum_{\text{Hydrogen bond}} \left( \frac{C_{sm}}{r_{sm}^{12}} - \frac{D_{sm}}{r_{sm}^{10}} \right) \\ &- \sum_{i,m} \frac{q_m}{r_{im}} + \sum_{k,m} \frac{Z_k q_m}{r_{km}} \\ &- \frac{1}{2} \sum_{i,m} \frac{\boldsymbol{\mu}_m \cdot \boldsymbol{r}_{im}}{r_{im}^3} + \frac{1}{2} \sum_{k,m} Z_k \frac{\boldsymbol{\mu}_m \cdot \boldsymbol{r}_{km}}{r_{km}^3} \end{split}$$

This approach will make  $H_{\rm QMMM}$  dependent on electronic coordinates, and thus  $H_{\rm QMMM}$  will not just be a number to be added to the total energy.

The terms in  $H_{\rm QMMM}$  involving electronic coordinates must be added to the HF SCF procedure, and in this case these terms in  $H_{\rm QMMM}$  will make the QM part feel the MM part also in the HF SCF procedure.

Look of the extra terms entering the Hartree Fock equations

Due to the double boxed expressions involving electronic coordinates the HF SCF procedure must be modified to include the resulting one electron contributions. The one electron integrals to be included in the HF SCF procedure are the following.

$$H_{lphaeta}^{\mathsf{QMMM}} = -\sum_m \left( \int \phi_lpha rac{q_m}{r_{im}} \phi_eta \; dr + rac{1}{2} \int \phi_lpha rac{oldsymbol{\mu}_m \cdot oldsymbol{r}_{im}}{r_{im}^3} \phi_eta \; dr 
ight)$$

As these extra terms are all one electron terms we may readily add them to the pure quantum mechanical Fock operator  $F_{\alpha\beta}^{\rm QM}$  thus producing another Fock operator  $F_{\alpha\beta}$  looking like the following.

$$F_{\alpha\beta} = F_{\alpha\beta}^{\mathsf{QM}} + H_{\alpha\beta}^{\mathsf{QMMM}}$$

Comment on inclusion of polarizability terms

It should be noted, that inclusion of polarizability terms in the HF SCF scheme induces a significant increase in computation time.

This is because the total restricted Hartree Fock wave function  $\Psi$  and the induced dipoles  $\mu$  are coupled and thus should be determined self consistently within the HF SCF scheme.

#### Approach I - The total energy

Assuming, that we are working in a closed shell system and thus applying a restricted Hartree Fock approach we end up with the following energy expression. Note, that  $E_{\rm MM}=H_{\rm MM}$  as well as  $E_{\rm QMMM}\underline{=}H_{\rm QMMM}$ .

$$E = E_{\rm QM} + E_{\rm MM} + E_{\rm QMMM}$$

$$E_{\rm QM} \ = \ \frac{1}{2} \sum_{\alpha\beta} P_{\alpha\beta} (H^{\rm QM}_{\alpha\beta} + F^{\rm QM}_{\alpha\beta}) + \sum_{k,l} \frac{Z_k Z_l}{r_{kl}}$$

$$\begin{split} E_{\text{MM}} &= \sum_{\text{bond}}^{\text{MM MM}} \frac{k_{\text{b}}}{2} (r - r_0)^2 + \sum_{\text{angle}}^{\text{MM MM}} \frac{k_{\text{a}}}{2} (\theta - \theta_0)^2 + \sum_{\text{dihedral}}^{\text{MM MM}} \frac{k_{\text{d}}}{2} (1 + \cos \Theta) \\ &+ \sum_{m,n} \left( \frac{A_{mn}}{r_{mn}^{12}} - \frac{B_{mn}}{r_{mn}^6} \right) + \sum_{\text{Hydrogen bond}} \left( \frac{C_{mn}}{r_{mn}^{12}} - \frac{D_{mn}}{r_{mn}^{10}} \right) \\ &+ \sum_{m,n} \frac{q_m q_n}{r_{mn}} \\ &+ \frac{1}{2} \sum_{m < n} q_m \frac{\boldsymbol{\mu}_n \cdot \boldsymbol{r}_{mn}}{r_{mn}^3} + \frac{1}{2} \sum_{m > n} q_m \frac{\boldsymbol{\mu}_n \cdot \boldsymbol{r}_{mn}}{r_{mn}^3} - \frac{1}{2} \sum_{m} \alpha_m \boldsymbol{E}_m \cdot \boldsymbol{E}_m \end{split}$$

$$\begin{split} E_{\text{QMMM}} &= \sum_{\text{bond}}^{\text{QM MM}} \frac{k_{\text{b}}}{2} (r - r_0)^2 + \sum_{\text{angle}}^{\text{QM MM}} \frac{k_{\text{a}}}{2} (\theta - \theta_0)^2 + \sum_{\text{dihedral}}^{\text{QM MM}} \frac{k_{\text{d}}}{2} (1 + \cos \Theta) \\ &+ \sum_{s,m} \left( \frac{A_{sm}}{r_{sm}^{12}} - \frac{B_{sm}}{r_{sm}^6} \right) + \sum_{\text{Hydrogen bond}} \left( \frac{C_{sm}}{r_{sm}^{12}} - \frac{D_{sm}}{r_{sm}^{10}} \right) \\ &+ \sum_{s,m} \frac{q_s q_m}{r_{sm}} \\ &+ \frac{1}{2} \sum_{s,m} q_s \frac{\boldsymbol{\mu}_m \cdot \boldsymbol{r}_{sm}}{r_{sm}^3} \end{split}$$

#### Approach II - The total energy

Assuming, that we are working in a closed shell system and thus applying a restricted Hartree Fock approach we end up with the following energy expression. Note, that  $E_{\rm MM}=H_{\rm MM}$  but  $E_{\rm QMMM} \neq H_{\rm QMMM}$ .

$$E = E_{\rm QM} + E_{\rm MM} + E_{\rm QMMM}$$

$$E_{\rm QM} \ = \ \frac{1}{2} \sum_{\alpha\beta} P_{\alpha\beta} (H^{\rm QM}_{\alpha\beta} + F^{\rm QM}_{\alpha\beta}) + \sum_{k,l} \frac{Z_k Z_l}{r_{kl}}$$

$$\begin{split} E_{\text{MM}} &= \sum_{\text{bond}}^{\text{MM MM}} \frac{k_{\text{b}}}{2} (r - r_0)^2 + \sum_{\text{angle}}^{\text{MM MM}} \frac{k_{\text{a}}}{2} (\theta - \theta_0)^2 + \sum_{\text{dihedral}}^{\text{MM MM}} \frac{k_{\text{d}}}{2} (1 + \cos \Theta) \\ &+ \sum_{m,n} \left( \frac{A_{mn}}{r_{mn}^{12}} - \frac{B_{mn}}{r_{mn}^6} \right) + \sum_{\text{Hydrogen bond}} \left( \frac{C_{mn}}{r_{mn}^{12}} - \frac{D_{mn}}{r_{mn}^{10}} \right) \\ &+ \sum_{m,n} \frac{q_m q_n}{r_{mn}} \\ &+ \frac{1}{2} \sum_{m < n} q_m \frac{\boldsymbol{\mu}_n \cdot \boldsymbol{r}_{mn}}{r_{mn}^3} + \frac{1}{2} \sum_{m > n} q_m \frac{\boldsymbol{\mu}_n \cdot \boldsymbol{r}_{mn}}{r_{mn}^3} - \frac{1}{2} \sum_{m} \alpha_m \boldsymbol{E}_m \cdot \boldsymbol{E}_m \end{split}$$

$$\begin{split} E_{\text{QMMM}} &= \sum_{\text{bond}}^{\text{QM MM}} \frac{k_{\text{b}}}{2} (r - r_0)^2 + \sum_{\text{angle}}^{\text{QM MM}} \frac{k_{\text{a}}}{2} (\theta - \theta_0)^2 + \sum_{\text{dihedral}}^{\text{QM MM}} \frac{k_{\text{d}}}{2} (1 + \cos \Theta) \\ &+ \sum_{s,m} \left( \frac{A_{sm}}{r_{sm}^{12}} - \frac{B_{sm}}{r_{sm}^6} \right) + \sum_{\text{Hydrogen bond}} \left( \frac{C_{sm}}{r_{sm}^{12}} - \frac{D_{sm}}{r_{sm}^{10}} \right) \\ &- \sum_{\alpha\beta} P_{\alpha\beta} \left( \sum_m \int \phi_\alpha \frac{q_m}{r_{im}} \phi_\beta \ dr \right) \right] + \sum_{k,m} \frac{Z_k q_m}{r_{km}} \end{split}$$

$$\boxed{ -\frac{1}{2} \sum_{\alpha\beta} P_{\alpha\beta} \left( \sum_{m} \int \phi_{\alpha} \frac{\boldsymbol{\mu}_{m} \cdot \boldsymbol{r}_{im}}{r_{im}^{3}} \phi_{\beta} dr \right) } + \frac{1}{2} \sum_{k,m} Z_{k} \frac{\boldsymbol{\mu}_{m} \cdot \boldsymbol{r}_{km}}{r_{km}^{3}}$$

### Example

QM $(6-31G*)$ and QMMM data from $[1]$			
$r_{C-CI}$ — Å	$\Delta E_{ m QM}$ — kcal/mole $^a$	$\Delta E_{ extsf{QMMM}}$ — kcal/mole $^b$	
6.014	0.0	0.0	
4.571	-2.6	-5.4	
3.274	-6.7	0.5	
2.682	-1.0	10.3	
2.383	7.2	18.3	

QM $(6-31G*)$ and QMSM data from [4]			
$r_{C-CI}$ — Å	$\Delta E_{QM}$ — kcal/mole $^c$	$\Delta E_{ extsf{QMSM}}$ — kcal/mole $^d$	
$\infty$	0.0		
7.500	-2.4	0.0	
6.000	-3.6		
4.500	-6.4		
3.269	-10.3		
2.974	-9.2		
2.678	-4.5		
2.530	-0.1		
2.383	3.6	28.0	

<sup>&</sup>lt;sup>a</sup>Energy differences scaled according to the  $r_{\rm C-Cl}=6.014$  structure <sup>b</sup>Energy differences scaled according to the  $r_{\rm C-Cl}=6.014$  structure <sup>c</sup>Energy differences scaled according to the  $r_{\rm C-Cl}=\infty$  structure <sup>d</sup>Energy differences scaled according to the  $r_{\rm C-Cl}=7.500$  structure

## Example

QM and QMMM data from [1]			
$r_{C-CI}$ — Å	$\Delta E_{ m QM}$ — kcal/mole $^a$	$\Delta E_{ extsf{QMMM}}$ — kcal/mole $^b$	
6.014	0.0	0.0	
4.571	-2.6	-5.4	
3.274	-6.7	0.5	
2.682	-1.0	10.3	
2.383	7.2	18.3	

$E_{QMMM}$ and $E_{QMMM}+E_{MM}$ data from $[1]$				
$r_{C-CI}$ — Å	$E_{ m QMMM}$ — kcal/mole	$E_{\mathrm{QMMM}} + E_{\mathrm{MM}}$ — kcal/mole		
6.014	-131.1	-1208.2		
4.571	-126.5	-1211.0		
3.274	-109.3	-1201.0		
2.682	-101.5	-1196.9		
2.383	-92.3	-1197.1		

 $<sup>^</sup>a \rm Energy$  differences scaled according to the  $r_{\rm C-Cl}=6.014$  structure  $^b \rm Energy$  differences scaled according to the  $r_{\rm C-Cl}=6.014$  structure

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$$- \ Lars \ Hemmingsen \left\{ \begin{array}{l} Royal \ Veterinary \\ and \\ Agricultural \ University \end{array} \right.$$

• Acknowledgements also go to all you listeners

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