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CORRECTION

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Correction: Benchmark and performance of longrange corrected time-dependent density functional tight binding (LC-TD-DFTB) on rhodopsins and light-harvesting complexes

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Correction for 'Benchmark and performance of long-range corrected time-dependent density functional tight binding (LC-TD-DFTB) on rhodopsins and light-harvesting complexes' by Beatrix M. Bold *et al.*, *Phys. Chem. Phys.*, 2020, **22**, 10500–10518, https://doi.org/10.1039/C9CP05753F.

The authors have recognised two errors in the data for the published version of this article. The first one concerns the reported QM/MM calculations on the Fenna–Matthews–Olson (FMO) complex, and the second error affects all Coulomb couplings presented.

In brief, the most important changes are: (i) the ZINDO site energies for the individual pigments are closer to each other and the ZINDO site energy fluctuations are smaller; (ii) the couplings are smaller and now in better agreement with reference data; and (iii) as a consequence, the exciton splitting is decreased. These errors do not change the conclusions of the study but only lead to small corrections of the reported results, as detailed below.

Error description

In the originally published version of the article, the excitation energies and couplings of the pigments in an FMO monomer were computed using the time-dependent long-range corrected density functional tight binding (TD-LC-DFTB) and ZINDO/S approaches in a QM/MM fashion representing their respective environment by point charges. The environment of an individual pigment consisted of the FMO monomer, the remaining six BChl a chromophores, TIP3P water molecules, as well as 15 sodium and 16 chloride ions.

Prior to the calculations of site energies and couplings along the trajectory, the coordinates of the system were processed as follows: the protein complex was centered in the simulation box with Gromacs tools in two steps. First, the protein and the chromophores were clustered using the *pbc cluster* option of *trjconv*; second, they were centered in the box using the *pbc mol* option of *trjconv*. However, it was not recognized that some counterions were placed outside the simulation box erroneously. Consequently, the MM point charges corresponding to these counterions had wrong coordinates in the QM/MM calculations of the site energies and couplings, leading to an incorrect description of the QM–MM electrostatics.

These calculations of site energies and couplings were repeated using a corrected procedure for the processing of the MM point charges. Specifically, the trajectory was centered using the Gromacs tools in a more appropriate way: in the first step, any appearing error due to the periodic boundary conditions were corrected, and the protein was centered in the box using the *pbc atom* option of *triconv* in the second step.

Additionally, the authors have recognized an error in a script that was used to compute the Coulomb couplings. The coordinates were not converted to atomic units as needed, which resulted in a significant overestimation of the coupling values reported in the original publication. The corrected couplings are now in much better agreement with the reference data.

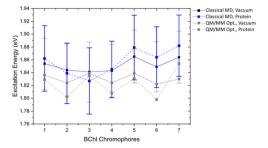
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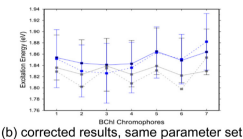
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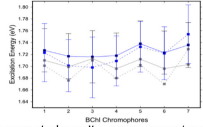
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shown in the published paper





(c) corrected results, new parameter set

Fig. 12 Excitation energies of the seven BChl a chromophores, computed for DFTB QM/MM optimized structures and sampled along the classical trajectories. These values were calculated in vacuum and with fixed MM point charges using LC-DFTB.

Corrections

All corrected results presented in this erratum were computed using the newest parameter set for excitation energies. In general, DFTB parameters can be divided into electronic parameters and parameters describing the repulsive potential. Excitation energy calculations as well as calculations of transition charges require only electronic parameters. These parameters can be optimized for certain properties with relatively low effort by variation of the compression radii. The parameters in the original publication were already optimized for excitation energies but the new parameter set derived with a small modification at the carbon compression radius provides improved absolute excitation energies. The predicted relative excitation energies of BChl a in FMO are basically the same, as shown in Fig. 12.

The authors recalculated only a selection and comment on the expected changes for the other figures and tables. In the following, they start with the benchmark of the couplings on the BChl a dimer models, followed by exemplary couplings in the LH2 complex. Thereafter, they continue with the new site energies, couplings and the resulting exciton energies of the FMO complex. Previous results are shown in parentheses, while the new results are shown in bold. The figures and tables carry the same numbers as in the published version of the article.

1 Benchmark: bacteriochlorophyll a (BChl a)-excitonic coupling I: model dimers

Table 3 shows that the correctly computed TD-LC-DFTB couplings not only have the same trend, but also the absolute values are very similar to the DFT references. Using the correction, the TD-LC-DFTB couplings are no more overestimated, but are slightly smaller than the reference values.

Table 3 Coulomb couplings (in eV) of the model system using the TrESP, Tr-Mulliken and LC-DFTB schemes. The partial charges of the TrESP and Tr-Mulliken values are either based on the B3LYP or the CAM-B3LYP functional

	Distance [Å]	7	8	9	10
TrESP	B3LYP	0.050	0.038	0.029	0.023
	CAM-B3LYP	0.056	0.043	0.033	0.026
Tr-Mulliken	B3LYP	0.047	0.036	0.028	0.022
	CAM-B3LYP	0.054	0.041	0.032	0.025
_	LC-DFTB	0.041 (0.082)	0.031 (0.063)	0.024 (0.049)	0.019 (0.039)

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Range of excitation and excitonic energies along a classical MD trajectory of the FMO complex. LC-DFTB is used for the computation of

excitation energies without (vacuum) and with the protein environment. The Coulomb couplings have been computed using LC-DFTB and TrESP. Moreover, the vacuum values have been slightly modified as well due to the newer parameter set applied in the present calculations

	Max	Min	Shift
Vacuum	1.738 (1.865)	1.716 (1.841)	-0.022 (-0.024)
Protein	1.754 (1.882)	1.698 (1.827)	-0.056 (-0.055)
Coupled chromophores	, ,	` ,	, ,
LC-DFTB	1.811 (1.979)	1.659 (1.775)	-0.152 (-0.204)
TrESP	1.876	1.777	-0.099
Experimental	1.563	1.503	-0.060

2 Performance of LC-DFTB in biological model systems: light-harvesting complexes: light-harvesting complex II (LH2)

The TD-LC-DFTB Coulomb couplings in LH2 are compared to the TrESP couplings in Fig. S5. The corrected TD-LC-DFTB couplings are now in much better agreement with the TrESP values. Within the B800 ring, they are slightly higher, but in the B850 ring the TD-LC-DFTB coupling values are of the same size as the TrESP couplings. The authors expect similarly smaller coupling values for all BChl a pairs in the LH2 complexes studied in the published version of the article.

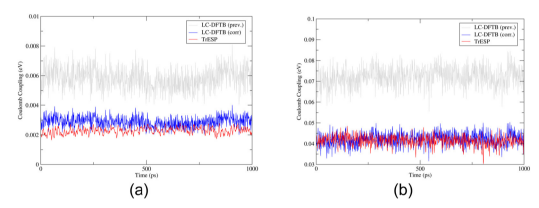


Fig. S5 Coulomb couplings sampled along a classical MD trajectory of 1 ns length for (a) one BChl a dimer pair with the strongest Coulomb couplings chosen as the example for the B800 ring and for (b) one BChl a dimer pair with the strongest Coulomb couplings chosen as the example for the B850 ring

Performance of LC-DFTB in biological model systems: light-harvesting complexes: Fenna-Matthews-Olson (FMO) complex

The erroneous electrostatics in the QM/MM calculations in the FMO complex had only a small effect on the TD-LC-DFTB site energies and couplings, while the ZINDO/S excitation energies show slightly larger deviations.

Fig. 12 depicts a slightly different trend for the corrected TD-LC-DFTB site energies compared to the ones shown in the original publication, while the range between the lowest and largest values remains basically the same (Table 9). The results in Fig. 12b

Table S19 Strongest Coulomb couplings (in eV) computed using LC-DFTB for the DFTB QM/MM optimized model of BChl a dimers and average Coulomb couplings as maxima of Gaussian fits of the MD sampled structures

BChl a dimer	QM/MM optimized	Classical MD
1-2	0.022 (0.044)	$0.017 \pm 0.003 \ (0.033 \pm 0.006)$
2-3	0.009 (0.019)	$0.008 \pm 0.001 \ (0.016 \pm 0.001)$
3-4	0.011 (0.021)	$0.012 \pm 0.003 \; (0.022 \pm 0.006)$
4-5	0.021 (0.043)	$0.021 \pm 0.003 \ (0.040 \pm 0.005)$
5-6	0.019 (0.037)	$0.015 \pm 0.003 \ (0.029 \pm 0.007)$
6-7	0.014 (0.028)	$0.014 \pm 0.003 \ (0.028 \pm 0.005)$
4-7	0.009 (0.019)	$0.009 \pm 0.003 \ (0.018 \pm 0.006)$

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Table S20 Maximum and minimum excitation and excitonic energies of the structures along an MD simulation of the FMO complex. ZINDO/S has been used for the computation of the excitation energies without (vacuum) and with the protein environment as fixed point charges. The Coulomb couplings have been computed using LC-DFTB and TrESP

	Max	Min	Shift
Vacuum	1.491	1.475	-0.016
Protein	1.496 (1.529)	1.465 (1.480)	-0.031 (-0.049)
Coupled chromophores	,	,	,
LC-DFTB	1.618 (1.657)	1.441 (1.436)	-0.177 (-0.221)
TrESP	1.648	1.446	-0.202
Experimental	1.563	1.503	-0.060
•			

were computed with the same parameter set as in the original publication and reveal the changes due to the corrected electrostatics. Fig. 12c shows the almost constant shift to lower excitation energies when the new parameter set is applied for the computations with a correct representation of the electrostatics.

The ZINDO/S approach now predicts a smaller range of the excitation energies along the MD trajectory (Fig. S7 and Table S20). It is remarkable that the excitation energy of BChl 5 is no longer overestimated as previously reported. Nonetheless, the large excitation energy of BChl 3 in the QM/MM optimized structure as well as the large energetic change in BChl 5 indicate a strong sensitivity to the electrostatics.

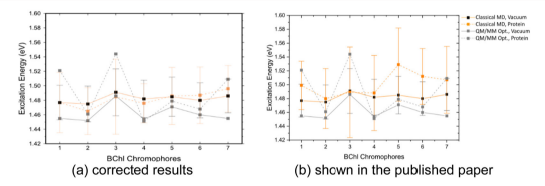


Fig. S7 Excitation energies of the seven BChl a chromophores, computed for DFTB QM/MM optimized structures and sampled along the classical trajectories. These values were calculated in vacuum and with fixed MM point charges using ZINDO/S.

The corrected couplings of the BChl a pairs in the FMO complex are presented in Table S19. Their values are roughly half of those previously presented in the published version of the article.

Finally, the excitonic energies were recalculated using the corrected excitation energy and coupling values (Table S21). The range of the excitonic energies is smaller due to the smaller couplings.

Table S21 Excitonic energies (in eV) of the FMO complex from C. tepidum based on the QM/MM optimized structures and averaged along a MD trajectory. The excitation energies and the Coulomb couplings have been computed using LC-DFTB

QM/MM optimized	MD simulation
1.656 (1.756)	1.659 (1.775)
1.662 (1.767)	1.686 (1.807)
1.670 (1.790)	1.707 (1.836)
1.707 (1.833)	1.726 (1.864)
1.714 (1.848)	1.749 (1.892)
1.721 (1.864)	1.774 (1.925)
1.721 (1.864)	1.774 (1.925)
1.738 (1.899)	1.811 (1.979)

The Royal Society of Chemistry apologises for these errors and any consequent inconvenience to authors and readers.

References

1 N. Schieschke, B. M. Bold, P. M. Dohmen, D. Wehl, M. Hoffmann, A. Dreuw, M. Elstner and S. Höfener, Geometry dependence of excitonic couplings and the consequences for configuration-space sampling, J. Comput. Chem., 2021, 42(20), 1402-1418.