Exciton $g$ factors of van der Waals heterostructures from first-principles calculations

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External fields are a powerful tool to probe optical excitations in a material. The linear energy shift of an exciton in a magnetic field is quantified by its effective $g$ factor. Here we show how exciton $g$ factors and their sign can be determined by converged first-principles calculations. We apply the method to monolayer excitons in semiconducting transition metal dichalcogenides and to interlayer excitons in MoSe$_2$/WSe$_2$ heterobilayers and obtain good agreement with recent experimental data. The precision of our method allows us to assign measured $g$ factors of optical peaks to specific transitions in the band structure and also to specific regions of the samples. This revealed the nature of various, previously measured interlayer exciton peaks. We further show that, due to specific optical selection rules, $g$ factors in van der Waals heterostructures are strongly spin- and stacking-dependent. The calculation of orbital angular momenta requires the summation over hundreds of factors of van der Waals heterostructures from first-principles calculations.

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I. INTRODUCTION

Since the dawn of quantum mechanics the application of external magnetic fields has proven to be an invaluable tool to probe the properties of matter. A good textbook example is the Zeeman effect in atoms, that describes the linear shift of an energy level $\varepsilon = g\mu_B B$ in a homogeneous magnetic field $B$, where $g$ is the Landé $g$ factor and $\mu_B$ is the Bohr magneton. The theory of magnetic field shifts in semiconductors is closely related and was developed by multiple authors before [1–3], mostly within the context of $k \cdot p$ perturbation theory or few-band tight-binding models. For conventional semiconductors, these models have proven to be useful and predictive but their applications to two-dimensional semiconductors based on transition metal dichalcogenides (TMD) has not led to satisfactory results yet [4–6]. Early experimental studies of the magnetic field dependence of excitons, i.e., optical excitations formed by bound electron-hole pairs, in monolayer MoSe$_2$ observed a Zeeman shift $g \approx -4$, which has been attributed to the $d$-orbital character of the conduction and valence states involved in the excitonic transition [7,8]. However, subsequent studies in WSe$_2$ [9–11] and WS$_2$ [12] where excitons exhibit the same orbital character, observed slightly larger values, which pointed to possible corrections due to the angular momentum texture of the conduction and valence bands. This picture became even more puzzling when $g$ factors of $\approx 9.5$ were experimentally observed for dark exciton states in bilayer WSe$_2$ [13], and when interlayer excitons in heterobilayers of TMD where demonstrated to have $g$ factors of $\approx 6.7$ and $\approx -16$ [14], which deviate even more from the value expected for ground state excitons in TMD. It is thus clear that a more rigorous theoretical model, which properly accounts for the angular momentum character of conduction and valence states in monolayer and bilayer materials, is required for a correct description of the exciton Zeeman shifts in these materials. In this work, we address this problem and offer a practical solution that particularly works for excitonic states.

To test and apply the method we consider monolayers (see Fig. 1) and heterobilayers (see Fig. 2) of TMD. They are particularly suited to our method because (i) their optical properties are dominated by excitons and (ii) related phenomena such as exciton complexes, Rydberg series, Zeeman shifts and more were recently studied in great detail [15,16].

A van der Waals heterostructure is formed by vertically stacking two-dimensional crystals via deposition or mechanical exfoliation. Today it is possible to fabricate heterostructures with arbitrary material sequence and relative lattice orientation (twist angle $\theta$) [17]. The interlayer interactions are weak and therefore many monolayer properties are preserved in heterostructures. TMD heterobilayers (HB) usually have a staggered (type-II) band alignment and free electrons and holes accumulate in different layers which leads to the formation of long-lived, charge-separated, spatially indirect interlayer excitons [18–20]. A mismatch of the in-plane lattice constants or a sufficiently large twist angle between individual layers leads to the formation of a moiré pattern where the lattice registry and the band gap continuously vary in space.

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FIG. 1. Properties of transition metal dichalcogenide monolayers MX\textsubscript{2}. (a) Top view of the atomic structure, large and small balls represent M (metal) and X (chalcogen) atoms, respectively. (b) The Brillouin zone with the points Γ at the center and K at the corners. The sign of the K points (valley index) alternates. (c) Schematic band structure at the +K point. Small arrows next to the colored bands indicate the spin orientation of the conduction (c, c + 1) and valence (v-1, v) bands. Double arrows indicate dipole-allowed optical transitions, where the polarization σ+ is shown in red, z in black and the dashed line represents a forbidden transition. In summary: the spin-conserving transitions at +K couple to σ+ polarized light, one of the spin-flip transitions is optically dark and the other one couples to z-polarized light.

This gap variation can act as an additional confining potential for interlayer excitons [21–23]. It was recently shown that in MoSe\textsubscript{2}/WSe\textsubscript{2} HB and MoS\textsubscript{2} bilayers with θ close to 0° (R) or 60° (H) structural deformations lead to strong deviations from the ideal moiré pattern and the areas of high-symmetry stacking configurations with the lowest total energies are significantly enlarged [24,25]. The period of these deformations is equal to the moiré wave length. For R systems the sample area is mostly covered by equal proportions of R\textsubscript{m} (AB) and R\textsubscript{h} (BA) stackings, while in H systems H\textsubscript{m} (ABBA) covers most of the sample [26]. This is illustrated in Figs. 2(a) and 2(b) where for the labeling of the stacking configurations [27] we follow the notation of Yu et al. [28,29].

In TMD monolayers, the fundamental band gap is direct and located at the corners of the hexagonal Brillouin zone at the ±K points [see Fig. 1(b)]. There are two symmetry inequivalent ±K valleys, that are connected by time-reversal symmetry, and the sign is called the valley index. Spin-orbit interactions split the band edge states into spin-polarized bands as indicated in Fig. 1(c). The magnitude of the splitting is several hundred meV in the valence band and only a few meV in the conduction band. Due to mirror symmetry in monolayers, the projection of the spin onto the quantization axis perpendicular to the layer is preserved and m\textsubscript{z} = 1/2 is a good quantum number. However, the spin-orbit coupling can lead to a reduction of m\textsubscript{z} while preserving m\textsubscript{z} = m\textsubscript{z} = 0, as shown in Ref. [30] for 2D hexagonal crystals. Nevertheless, in most cases, taking m\textsubscript{z} = 1/2 was demonstrated to be a reasonable approximation [31]. In molybdenum-based monolayers the spin orientation of the valence and conduction bands is the same, while in tungsten-based systems the spin orientation is opposite [32]. At the ±K valleys optical transitions couple to light of specific circular (σ±) or linear (z) polarization, as indicated by vertical double arrows in Fig. 1(c). The allowed transitions are determined by dipole selection rules,

\[ |\mathbf{e}_+ \cdot \mathbf{\pi}_{\text{c}(k)}|^2 > 0 \iff \sigma^+, \]
\[ |\mathbf{e}_- \cdot \mathbf{\pi}_{\text{c}(k)}|^2 > 0 \iff \sigma^-, \]
\[ |\mathbf{z} \cdot \mathbf{\pi}_{\text{c}(k)}|^2 > 0 \iff \mathbf{z}, \]

where \( \mathbf{e}_\pm = (1, \pm i, 0)/\sqrt{2} \), \( \mathbf{z} = (0, 0, 1) \), and \( \mathbf{\pi}_{\text{c}(k)} = (\pi_{\text{c}(k)}, \pi_{\text{c}(k)}, \pi_{\text{c}(k)}^*) \) are momentum (or optical) matrix elements for transitions between the valence and conduction band and \( \nu, \sigma \) are the corresponding band indices. The left-hand side of Eq. (1) is directly proportional to the oscillator strength of a transition and therefore we will refer to it as “intensity.” The selection rules differ in monolayers and HB, where they are also stacking-dependent [29]. In Fig. 1(c) it is discernible that in monolayers the spin-conserving transition (giving rise to spin-singlet excitons) couples to σ+ light at the +K valley (and to σ− at −K) and one spin-flip transition (leading to spin-triplet excitons) couples to z-polarized light and the other one is forbidden/dark. In stark contrast are the selection rules of MoSe\textsubscript{2}/WSe\textsubscript{2} HB, that are shown in Fig. 2(c). There, depending on the stacking configuration, spin-conserving and spin-flip transitions couple to entirely different polarizations, e.g., for the spin-conserving transition in a R\textsubscript{h} HB we have (σ± ↔ ±K), while in a R\textsubscript{h} HB we have (σ± ↔ ±K).

In this paper we demonstrate how the theory of magnetic field-induced energy shifts in semiconductors can be realized with state of the art density functional theory calculations. We test the method by calculating g factors of excitons in MoS\textsubscript{2}, MoSe\textsubscript{2}, MoTe\textsubscript{2}, WS\textsubscript{2}, WSe\textsubscript{2} monolayers and obtain excellent agreement with available experimental data. Then, we consider interlayer excitons in MoSe\textsubscript{2}/WSe\textsubscript{2} HB (which might serve as model for arbitrary TMD-based HB) and show that the approach can explain recent magnetooptical
measurements on HB, where unusual signs and values of excitonic $g$ factors were reported [14,33–35]. We further demonstrate how stacking-dependent selection rules lead to stacking dependent exciton $g$ factors.

II. THEORY OF MAGNETIC FIELD SHIFTS IN SEMICONDUCTORS

A. Effective $g$ factor of a Bloch state

The basic theory of the magnetic field dependence of Bloch states has been developed before by multiple authors and is usually applied in models [1–6,36]. Here we reformulate it in a way suitable for general electronic structure calculations. The starting point is a nonrelativistic band structure Hamiltonian $H^0$ and its corresponding band energies $\varepsilon_{nk}^0$, and Bloch states $|nk\rangle$ (i.e., Bloch phase times lattice-periodic function),

$$H^0 = \frac{p^2}{2m_0} + V,$$

$$H^0 |nk\rangle = \varepsilon_{nk}^0 |nk\rangle,$$

$$1 = \sum_n |nk\rangle \langle nk|,$$

where $p$ is the momentum operator, $m_0$ is the rest mass of the electron, $V$ is the effective potential, and $n$ and $k$ are the band index and the wave number, respectively. The last line emphasizes that the set of Bloch states forms a complete basis. These states are obtained from electronic structure calculations and are supposed to be known. The coupling of these states to an external magnetic field is described by adding the spin Zeeman term to $H^0$ and by replacing the momentum operator $p$ by $p - qA$ (minimal coupling), where $A$ is the vector potential, $q = -|e|\hbar$ the charge of the electron and $e_0$ is the elementary charge. For a uniform external magnetic field $B$ it is convenient to choose $A = (B \times r)/2$, which satisfies the Coulomb gauge $\nabla \cdot A = 0$, where $r$ is the position operator. This leads to the Pauli equation

$$H(B) = H^0 + H^1(B) + H^Q(B)$$

$$= H^0 + \mu_B B \cdot (L + \frac{g_0}{2} \Sigma) + \frac{e_0}{8m_0} (B \times r)^2,$$

with the matrix elements $L_{nk} = \langle nk| L^z |nk\rangle$, $\Sigma_{nk} = \langle nk| \Sigma^\pm |nk\rangle$ and $H^Q(B) = e_0 B^2/8m_0 (|r|^2 + (r')^2) |nk\rangle$. The effective $g$ factor of the Bloch state $|nk\rangle$ is thus

$$g_{nk} = L_{nk} + \Sigma_{nk}.$$

The orbital angular momentum matrix elements are evaluated as

$$L_{nk} = \frac{1}{\hbar} \langle nk| r^x p^y - r^y p^x |nk\rangle$$

$$= \frac{1}{\hbar} \sum_{m=1}^N \int r_m n_m p_{m,nk} - r_{m,nk} p_{m,nk}$$

$$= \frac{1}{\hbar} \sum_{m=1}^N n_m p_{m,nk} - p_{m,nk}^*,$$

Now choosing $B = (0, 0, B)$ parallel to the Cartesian $z$ direction and $g_0/2 \approx 1$ we get

$$\varepsilon_{nk}(B) = \varepsilon_{nk}^0 + \mu_B B (L_{nk} + \Sigma_{nk}) + H^Q_{nk},$$

with the matrix elements $L_{nk} = \langle nk| L^z |nk\rangle$, $\Sigma_{nk} = \langle nk| \Sigma^\pm |nk\rangle$ and $H^Q_{nk} = e_0 B^2/8m_0 (|r|^2 + (r')^2) |nk\rangle$. The effective $g$ factor of the Bloch state $|nk\rangle$ is thus

$$g_{nk} = L_{nk} + \Sigma_{nk}.$$

Equation (8) can be applied not only to Bloch states of crystals, but also to atoms or molecules. For an atomic orbital (ao) of the hydrogen atom it can be shown that for a sufficiently large number of states $N$, included in the summation, this expression converges to the well known analytical result $L_{ao} = m^2/2m = (n'l^m'|L_n)|n'l^m'\rangle = m^2/2m$. However, the convergence is slow. In the literature on TMD $L_{nk}$ is sometimes divided into a contribution coming from the atomic orbital (ao) and one from the lattice (l) (or valley) $L_{nk} = L_{ao} + L_{l,nk}$ and the two contributions are separately discussed [9,11,14,33]. However, this division is only of qualitative nature, as the projection of a Bloch state $|nk\rangle$ onto atomiclike orbitals is nonunique and leads to contributions from multiple atomiclike orbitals.
1. Relativistic effects

Above, we outlined the nonrelativistic theory that is satisfactory for light elements, but for systems with heavier atoms (such as Mo and W) relativistic effects cannot be neglected. In this paper we are mostly concerned with electronic structure calculations based on density functional theory (DFT). Relativistic effects and external magnetic fields can be introduced into DFT via current density functional theory [41,42].

However, for valence states it is sufficient to consider a 2-spinor formulation for an approximate relativistic Hamiltonian $H^{0,\text{rel}} = H^0(p^2) + H^{\text{SOC}}(p) + H^{\text{MV}}(p^4) + H^0 + mc^2$, where $H^0$ is Hamiltonian (2) and the other terms represent the spin-orbit coupling, the mass-velocity relation, the Darwin shift and the electron rest mass, respectively [43]. Neglecting the spin-orbit term leads to a scalar-relativistic approach, that is often used in solid state codes [44].

In g-factor calculations including relativistic effects $H^0$ in Eq. (2) is replaced by $H^{0,\text{rel}}$ which defines the set of unperturbed Bloch states. Then the coupling of $H^{0,\text{rel}}$ to the magnetic field is realized by adding the spin Zeeman term and replacing $p$ by $p - g\mathbf{A}$ in the parts that explicitly depend on $p$. For $H^{0}(p^2)$ this procedure leads to Eq. (5). In TMD systems the coupling of $H^{\text{MV}}(p^4)$ leads to marginal corrections that are neglected here. This leaves $H^{\text{SOC}}(p)$, which gives an additional linear contribution that is taken into account by replacing the momentum operator $p$ in $H^1(\mathbf{B})$ by

$$\mathbf{p} = \mathbf{p} + \frac{\hbar}{4m_0c^2} \sum \mathbf{\nabla} \Phi.$$  

(9)

Specifically, $\pi^a_{\text{m} \text{n} \text{k}}$ needs to be replaced by $\pi^a_{\text{m} \text{n} \text{k}} = \langle n\mathbf{k}|\pi^a|m\mathbf{k}\rangle$ in Eq. (8). Mind that this replacement also affects the optical selection rules [see Eq. (1)], where spin-orbit coupling enables spin-flip transitions.

B. Effective g factor of excitons

Excitons are bound states formed by electron and holes from the conduction (c) and valence (v) band edges, respectively. Using Eq. (6) we define the momentum-direct exciton energy as

$$E_{\mathbf{k}}(\mathbf{B}) = \epsilon_{\mathbf{k}c}(\mathbf{B}) - \epsilon_{\mathbf{k}v}(\mathbf{B}) - E_{\mathbf{k}}^{\text{Binding}} = E_{\mathbf{k}}^0 + E_{\mathbf{k}}^{\text{L}}(\mathbf{B}) + E_{\mathbf{k}}^{\text{Q}}(\mathbf{B}),$$  

(10)

where $E_{\mathbf{k}}^{\text{Binding}}$ is the exciton binding energy (that varies throughout the Brillouin zone), $E_{\mathbf{k}}^0 = \epsilon_{\mathbf{k}c} - \epsilon_{\mathbf{k}v} - E_{\mathbf{k}}^{\text{Binding}}$ is the zero-field exciton energy, $E_{\mathbf{k}}^{\text{Q}}(\mathbf{B}) = H_{\mathbf{k}}^{\text{Q}} - H_{\mathbf{k}}^{\text{v}}$ is the quadratic shift. The linear shift is

$$E_{\mathbf{k}}^{\text{L}}(\mathbf{B}) = (g_{\mathbf{k}c} - g_{\mathbf{k}v})\mu_0B = g_{\mathbf{k}}\mu_0B$$  

(11)

and $g_{\mathbf{k}}$ is the intravalley g factor of an exciton at $\mathbf{k}$.

It is also possible to consider momentum-indirect excitons, where electron and hole originate from Bloch states with different crystal momentum $\mathbf{k}$ [45].

III. NUMERICAL METHODS

The electronic structure calculations were performed with density functional theory (DFT) using the Vienna ab initio simulation package (VASP) [46] version 5.4.4, Perdew-Burke-Ernzerhof (PBE) [47] exchange-correlation functional and the Projector Augmented Wave method [48] with potentials of version 54. For testing purposes, we also used the local density approximation (LDA). An energy cutoff of 300 eV and a $6 \times 6 \times 1$ k mesh were chosen after careful convergence tests. The $k$-space integration was carried out with a Gaussian smearing method using an energy width of 0.05 eV for all calculations. All unit cells were built with at least 15 Å separation between replicates in the perpendicular direction to achieve negligible interaction. Dispersion interactions corrections were of Tkachenko-Scheffler (TS) type [49]. Atomic positions and lattice constants were optimized with $10^{-3}$ eV/Å and 0.1 kbar precision. The optimized values are given in footnote [50]. A comparative calculation for WS$_2$ was performed with the all-electron, full-potential linearised augmented plane wave (LAPW) method as implemented in the ELK package, using default parameters [51]. The momentum matrix elements $\pi_{\text{m} \text{n} \text{k}}^{\sigma}$ in VASP were obtained from the wave function derivatives that are calculated within density functional perturbation theory [52], in ELK they were calculated according to Eq. (9).

IV. RESULTS AND DISCUSSION

A. Transition metal dichalcogenide systems and the impact of optical selection rules on g factors

In TMD monolayers and heterostructures the band edge states are mostly at $\mathbf{k} = \pm \mathbf{K}$, which is what we will focus on in this article. Due to time-reversal symmetry $\Sigma_{\mathbf{n} \mathbf{K}}^{\pm} = -\Sigma_{\mathbf{n} \mathbf{-K}}^{\pm}$ and $L_{\mathbf{n} \mathbf{+K}} = -L_{\mathbf{n} \mathbf{-K}}$. Spin-orbit interactions split the band edge states of monolayers into spin-polarized bands [see Fig. 1(c)] and $\Sigma_{\mathbf{v} \mathbf{\pm K}} = \pm 1$ is commonly assumed [16]. We use this specific property to define the valley index; so the valley where the valence band maximum is spin-up is $\mathbf{+K}$. In fact, $ab$ initio calculations of monolayer TMD show that $|\Sigma_{\mathbf{n} \mathbf{\pm K}}| < 1$ at the band edge ($n = u, v - 1, c, c + 1$). However, the effect is so small that it has a negligible influence on the g factor [53]. In TMD HB such calculations also show highly spin polarized band edge states at the K points [54]. Therefore, taking $\Sigma_{\mathbf{n} \mathbf{\pm K}} = \pm 1$ for those states is indeed a reasonable approximation. For the Bloch state and exciton g factors the above symmetry properties imply $g_{\mathbf{n} \mathbf{+K}} = -g_{\mathbf{n} \mathbf{-K}}$ and $g_{\mathbf{+K}} = -g_{\mathbf{-K}}$, respectively.

The valley-dependent selection rules, as discussed in the Introduction and visualized in Figs. 1 and 2, are employed to experimentally determine the excitonic g factors, where it is common to use

$$E_{\mathbf{\sigma} \mathbf{+K}} - E_{\mathbf{\sigma} \mathbf{-K}} = g_{\mathbf{\mu_0B}}$$  

(12)

to extract the linear magnetic shift and to define the intervalley g factor $g$. Using Eqs. (10) and (11) it follows for the lowest energy transition in MoS$_2$ monolayers (A exciton) $g_{\mathbf{A}}^{\mathbf{\text{L}}} = g_{\mathbf{+K}} - g_{\mathbf{-K}} = g_{\mathbf{+K}} - g_{\mathbf{-K}} = 2g_{\mathbf{+K}}$. In a $R^4$ HB the selection rules are the same and we obtain the same result $g_{\mathbf{R}^4}^{\mathbf{\text{L}}} = 2g_{\mathbf{+K}}$. But a $R^4$ HB has different selection rules and therefore $g_{\mathbf{R}^4}^{\mathbf{\text{L}}} = g_{\mathbf{+K}} - g_{\mathbf{-K}} = g_{\mathbf{+K}} - g_{\mathbf{-K}} = 2g_{\mathbf{-K}}$. This demonstrates that in HB the sign of the intervalley g factors, as defined by Eq. (12),
depends on the stacking configuration, which will further be discussed below.

B. Exciton g factors of monolayers

To apply this first-principles approach, we first consider TMD monolayers since they are well-studied and therefore represent a good test case. However, previous attempts to calculate the $g$ factor of TMD monolayers without making assumptions about the orbital moment contributions were not very satisfactory [4-6] — a problem that the present approach can solve. For the $g$ factors of A and B excitons Eqs. (12), (11), and (7) give $g_{A,B}^{II} = 2g_{v+K} = 2(\Delta \Sigma_{v+K} + \Delta L_{v+K})$, where $\Delta \Sigma_{v+K}$ and $\Delta L_{v+K}$ are the difference of the spin and the orbital angular momentum expectation values between conduction and valence band, respectively. Figure 1(c) shows that circular polarized light couples valence and conduction band states with the same spin, consequently $\Delta \Sigma_{v+K} = 0$ and only $\Delta L_{v+K}$ matters. In WS$_2$ the A (B) excitons are formed by the transitions $\nu \rightarrow c + 1$ ($\nu - 1 \rightarrow c$) and therefore $g_{A,B}^{II} = 2(L_{\nu+1,v+K} - L_{\nu-1,v+K})$ and $g_{A,B}^{II} = 2(L_{\nu+1,v+K} - L_{\nu-1,v+K})$.

Figures 3(a) and 3(b) show the convergence of $L_{v+K}$ and $g_{v+K}$ with respect to the number of bands $N$ included in the calculation [Eq. (8)] for WS$_2$. The convergence behavior of the other considered TMD is shown in Fig. 4. The largest contribution to $L_{v+K}$ is at the band gap (dashed vertical line) because the energy denominator in Eq. (8) is smallest there, but apart from that, the convergence is very slow. We find that for all considered TMD and the PBE-PAW method around $N = 300-500$ states are required to converge both quantities to a precision of 0.1 and around 700-900 to obtain an accuracy of 0.01 (for details see Fig. 4). The slow convergence can be understood by noticing that TMD monolayers strongly absorb light over a broad energy range [69], which means that there are many optical transitions with high intensities (momentum matrix elements) that contribute to Eq. (8). This slow convergence is in contrast to conventional semiconductors, where only a few bands are required to obtain convergence [2]. This finally explains why previous attempts to calculate exciton $g$ factors with few-band models did not lead to satisfactory results [4-6] — the orbital contributions were not converged.

Figure 3 also shows that for the same geometry the PBE and LDA results, obtained with the plane-wave-based, frozen-core PAW method (PBE-PAW and LDA-PAW) and the all-electron, full-potential LAPW method (PBE-LAPW) are nearly identical. This shows that our results are consistent and not bound to a specific code or (semi)local functional; the small differences are due to numerical reasons.

It is well-known that standard DFT calculations using (semi)local functionals like PBE or LDA underestimate band gaps. This overestimates $L_{v+K}$, due to the energy denominator in Eq. (8). Quasiparticle GW calculations are able to correct this error but they are numerically expensive. Fortunately the wavefunctions obtained from (semi)local DFT are almost identical to GW wavefunctions [70,71] (which explains why non-self-consistent approaches like $G_0W_0$ give reasonable results). Therefore, we expect the DFT spin and momentum matrix elements $\Sigma_{\nu\nu\nu}$ and $\pi_{\nu\nu\nu}$ to be reasonable and it is a good approximation to only correct the eigenvalue spectrum, in particular the band gaps. This is conveniently done by
defining a “scissor operator,”

\[ \varepsilon_{\text{sc}}' = \begin{cases} 
\varepsilon_{\text{sc}} + \Delta, \\
\varepsilon_{\text{sc}} 
\end{cases} \]

that modifies the band energies by simply increasing the band gap by \( \Delta \). As shown in Fig. 3(c), \( L_{m,+K} \) decreases with \( \Delta \). When increasing the band gaps of the considered TMD to their \( G_0 W_0 \) value [72] (see dashed vertical line) the \( L_{m,+K} \) decrease by values ranging from 0.50–1.13 (17–44%). These are big changes, which shows that calculating \( L_{m,+K} \) for individual bands is challenging. The individual \( g \) factors of conduction or valence bands could be probed separately via transport experiments and this could provide some insight to identify the individual values. However, the changes of the conduction and valence band states are very similar and when taking their change for calculating the exciton \( g \) factor, the band gap dependence nearly disappears. This is discernible in Fig. 3(d); \( g_{A,B}^{1L} \) of TMD increase only by 0.15–0.18 (3.9–4.7%) when the band gap is increased to the \( G_0 W_0 \) value. These changes are small enough to claim that standard DFT calculations using semilocal functional are suitable for calculating exciton \( g \) factors. Therefore, we do not apply the “scissor operator” to the results below.

In Table I we provide the PBE-PAW \( g \) factors for the considered TMD, which are approximately equal to \(-4\) for all systems. The experimental values, provided in the table, have a quite large statistical spread, even when we limit ourselves to undoped, encapsulated samples and measurements at \( T = 4 \) K. However, all values are negative and vary about \(-4\), which is fully consistent with our theoretical results. To our knowledge, this represents the first successful, parameter-free calculation of exciton \( g \) factors in TMD. Overall, we do not find significant differences in the \( g \) factors and the orbital angular momenta between the TMD monolayers. However, the calculated intensities in \( W_{X_2} \) are larger than in \( MoX_2 \) systems, which is consistent with measured photoluminescence spectra at room temperature [15]. The orbital angular momenta at \(+K\) in Table I are all positive and much bigger than commonly assumed in the literature, where \( L \) is often approximated by the atomic orbital contribution \( L_{ao}^{m,+K} = L_{e,+K}^{ao} \). However, \( \Delta L \) is always close to \(-2\), which explains the success of these simple models. In Table I the \( g \) factors of both \( A \) and \( B \) excitons are given. The two values are quite similar and they are close to \(-4\) in all systems. But we consistently find that \( g_{A}^{1L} > g_{B}^{1L} \), which agrees with some experimental findings [57,64].

**C. Stacking- and spin-dependent \( g \) factors of interlayer excitons in heterobilayers**

Now we apply the method to interlayer excitons in van der Waals heterostructures. As prototypical moiré system we chose \( MoSe_2/WSe_2 \) HB where unexpected values of \( g \) factors were recently reported [14,33–35]. The lattice constants of the monolayers are almost identical and for precise twist angles of \( \theta \approx 0^\circ \) (R) or \( 60^\circ \) (H) (and multiples of it) the system is (quasi) commensurate [74]. But when samples are fabricated by exfoliation methods \( \theta \) cannot be precisely controlled; for \( \theta \approx 0^\circ \) or \( \theta \approx 60^\circ \) the lattice reconstructs and certain high-symmetry stacking configurations dominate the sample [see Fig. 2(a)] [24,25]. Thus, it is sufficient to only study those high-symmetry stacking configurations, because they represent most of the properties of the HB.

The calculated \( g \) factors of K point interlayer excitons for each of these stackings are given in Table II. These values show explicitly that \( g \) factors in TMD HB are spin- and stacking-dependent, as discussed in Sec. IV A. Also indicated are the corresponding optical transitions between the valence \((v)\) and the conduction \((c, c + 1)\) bands and their intensities, which are two to three orders of magnitude lower than the ones of monolayer transitions (see Table I). This agrees well with previous results [29,75,76] and explains why interlayer excitons are hard to observe by absorption spectroscopy and are typically probed in photoluminescence experiments. The intensities of \( R^M_h \) and \( H^M_h \) are significantly lower and the
TABLE II. Calculated g factors $g^\text{HB}$ [Eq. (12)] of interlayer excitons for high-symmetry stacking configurations of MoSe$_2$/WSe$_2$ heterobilayers and comparison with reported experimental values. Also indicated are the corresponding transitions between the valence band (v) and the conduction (c, c + 1) band at the +K point, their intensities $\frac{\Delta_0}{\Sigma_0} (e \cdot \pi)^2$ in (eV Å)$^2$, circular polarizations and whether it is a spin-conserving ($\uparrow \uparrow$) or a spin-flip ($\uparrow \downarrow$) transition. $\Delta_\Sigma = \Sigma_{c + K} - \Sigma_{c - K}$ is the spin contribution (where $|\Sigma_{c + K}| = 1$ is used) and $\Delta L = L_{c + K} - L_{c - K}$ is the orbital contribution to $g^\text{HB}; L_0 = L_{c + K}$. All results are obtained with the PBE-PAW method. The $g$ factors are strongly stacking-dependent. Good agreement with experiment is found for v $\rightarrow$ c transitions with sizable intensities (highlighted).

| Transition          | $R_h^x$ | $R_h^y$ | $R_h^\Sigma$ | $H_h^x$ | $H_h^y$ | $H_h^\Sigma$ |
|---------------------|---------|---------|--------------|---------|---------|------------|
| v $\rightarrow$ c   | 6.19    | -10.73  | -6.15        | 10.42   | -12.60  | -16.67     |
| v $\rightarrow$ c+1 | 6.72    | -10.6$^b$ | -8.5$^b$    | 15.89$^c$ | -15.1$^c$ | 10.7$^d$  |
| v $\rightarrow$ c+1 | 6.99$^e$ |         |              |         |         |            |
| Spin                | $\uparrow \uparrow$ | $\uparrow \downarrow$ | $\uparrow \uparrow$ | $\uparrow \downarrow$ | $\uparrow \downarrow$ | $\uparrow \downarrow$ |
| Intensity           | 0.08    | 0.05    | 0.12         | 10$^{-7}$ | 0.01    | 0.03       |
| Polarization        | $\sigma -$ | $\sigma +$ | $\sigma -$ | $\sigma +$ | $\sigma -$ | $\sigma -$ |
| Spin                | $\uparrow \downarrow$ | $\uparrow \uparrow$ | $\uparrow \uparrow$ | $\uparrow \downarrow$ | $\uparrow \downarrow$ | $\uparrow \downarrow$ |
| $\Delta \Sigma$     | 1.80    | 1.53    | 1.79         | 1.53    | -1.53   | -1.79      |
| $L_{c+1}$           | 4.90    | 4.90    | 4.86         | 4.74    | 4.77    | 4.54       |
| $\Delta L$          | -3.10   | -3.37   | -3.08        | -3.21   | -6.30   | -6.34      |

$^a$Reference [14]; $^b$Reference [34]; $^c$Reference [33]; $^d$Ref. [35], the authors only measured $|g^\text{HB}|$; $^e$Ref. [73]; $^f$Reference [73], value of charged exciton.

transitions can probably not be observed. If we further consider that experimentally $g$ factors are determined by low-temperature photoluminescence spectroscopy where only the lowest energy transition (v $\rightarrow$ c) matters, then we are left with interlayer exciton $g$ factors of +6.2, and $-$6.2 for 0° (R) and $-$16.7 for 60° (H) systems (highlighted in Table II). Taking into account the large statistical spread of reported experimental $g$ factors (see Table I), these values are almost in quantitative agreement with recent experiments on MoSe$_2$/WSe$_2$ HB, where we are able to match our result to measurements of Seyler et al. (6.72 ± 0.02 for $\theta \approx 2^\circ$ and $-$15.89 ± 0.03 for $\theta \approx 57^\circ$), Ciarrocchi et al. (+7.1 ± 1.6 and $-$8.5 ± 1.5 for $|\theta| < 1^\circ$), Nagler et al. ($-$15.1 ± 0.1 for $\theta \approx 54^\circ$), and Joe et al. ($+$6.99 ± 0.35) [14,33,34,73]. The $g$ factor of $-$16.7 originates from regions with H$_h^\Sigma$ stacking, which is also covering most of the sample [see Fig. 2(a)]. In R systems $g = +6.2$ is linked to R$_h^x$, which is the dominant stacking (together with R$_h^y$). The negative $g$ factor $-$6.2 comes from regions with R$_h^y$ stacking, that is present only in small parts of the samples (the nodes). Ciarrocchi et al. [34] ascribe their $g = -8.5$ peak to the spin-conserving and the $g = +7.1$ peak to the spin-flip transition of the R$_h^x$ stacking. However, in Table II the signs of the calculated $g$ factors of spin-conserving and spin-flip transitions of R$_h^x$ are exactly opposite to their interpretation and the magnitudes of these two $g$ factors differ substantially. Therefore, our results suggest that the two peaks reported by Ciarrocchi et al. are related to spin-conserving transitions and they originate from different parts of the sample. For H systems Wang et al. [35] find two transitions with $g$ factor magnitudes of $|g^\text{HB}| = 15.2 \pm 0.2$ and $10.7 \pm 0.2$ (it is important to note that the authors did not determine the sign of their $g$ factors) and assign them to spin-singlet and spin-triplet excitons, which correspond to spin-conserving and spin-flip transitions, respectively. Our first-principles results give slightly bigger magnitudes but otherwise confirm this assignment. The spin-conserving transitions of H$_h^x$ and H$_h^\Sigma$ are both candidates to explain the lower of the two values; still it is more likely that the transition originates from H$_h^\Sigma$ because the samples are mostly covered by H$_h^\Sigma$ stackings and the oscillator strength of the transition is particularly large. In electron-doped R-type samples Joe et al. measure a PL peak with $g^R = -10.6 \pm 1.0$ and in undoped samples they find $+6.99 \pm 0.35$ [73]. The authors ascribe these two peaks to charged and neutral interlayer excitons, respectively. According to Wang et al. the approach for calculating $g$ factors of neutral and charged excitons is the same [5]. Our values of $-10.7$ for the spin-flip (v $\rightarrow$ c + 1) transition and +6.2 for the spin-conserving transition (v $\rightarrow$ c) transition in the R$_h^x$ stacking nicely agree with these measurements. However, more detailed analysis will be necessary to fully understand the agreement for charged excitons. The remaining predicted values we present in Table II could be observed in future experiments.

After showing the good agreement with recent experiments, let us now analyze orbital and spin contributions and the sign of the $g$ factors. In MoSe$_2$/WSe$_2$ HB the band alignment is such that MoSe$_2$ states form the conduction band and WSe$_2$ states the valence band. This is indicated by the color code in Figs. 2(b) and 2(c). In TMD HB the K-point states do not hybridize and are basically a superposition of monolayer states [77]. That is why the magnitudes of L$_h$ in Table II deviate only marginally from the corresponding monolayer values. In H systems the real space twist of the monolayers relative to each other is connected to a similar twist of the Brillouin zones. Hence for H systems the MoSe$_2$ conduction band state from $-K$ is at $-15.2 \pm 1$ for $\theta \approx 54^\circ$. This swaps the sign of the related spin and orbital contributions, as presented in Table II by the negative value of L$_{c+1}$ for H systems. As a consequence the orbital contribution $\Delta L$ of H systems is approximately twice the value of R systems, which explains why the magnitude of the $g$ factors is always bigger for H than for R systems. In HB both spin-conserving ($\uparrow \uparrow$) and spin-flip transition ($\uparrow \downarrow$) can couple to circularly
polarized light and hence they matter when defining the $g$ factor via Eq. (12). Furthermore a spin-flip transition provides a spin contribution to the $g$ factor of $\Delta \Sigma = -2$, that generally increases the magnitude of the $g$ factor. This is most significant for the $g$ factor of $-16.7$ for the H$_2^1$ stacking configuration. The large magnitude is a consequence of it being (i) a H transition and (ii) a spin-flip transition (leading to a spin-triplet exciton) [35]. If we consider the intra-valley $g$ factor at $+K$, as defined by Eq. (11), then all $g$ factors would be negative, because only $\Delta L$ and $\Delta \Sigma$ matter. However, the intervalley $g$ factor, according to Eq. (12) and commonly used in experiment, employs valley selection rules for circularly polarized light. The stacking- and spin-dependence of these selection rules is what leads to $g$ factors with both positive and negative signs. For example, the intravalley $g$ factor at $+K$ for the R$_h^1$ stacking is $g_{+K} = \Delta L + \Delta \Sigma = -3.10$ and $g_{-K} = +3.10$, due to time-reversal symmetry. Then applying the corresponding optical selection rules to obtain the intervalley $g$ factor gives $g_{\Delta L} = g_{+\Delta L} - g_{-\Delta L} = g_{+K} - g_{-K} = +6.2$. In many HB samples multiple interlayer exciton peaks are experimentally found and not all of them can be explained by considering momentum direct K-point transitions. It is likely that momentum-indirect excitons are playing an important role in these systems [45].

Let us now have a look at the electron $g$ factor. Jian et al. reported a value of $+1.07 \pm 0.079$ at $+K$ (and $-1.11 \pm 0.095$ at $-K$) but they were not able to determine if their sample is R or H [78]. Using the results in Table II and Eq. (7) we obtain $g_{c,+K} = +2.8$ for R stackings and the same value with negative sign for H stackings. Considering that the orbital contribution is calculated without scissor correction, we expect the actual $g$ factor to be smaller. If we now assume that the sign convention of Jian et al. is consistent with ours, then our results indicate that their system is of R type (i.e., $\theta \approx 0$). Thus, $g$ factor measurements of excitons (or even electrons) combined with our results enable to determine whether a system is R or H. For exfoliated HB such a tool is sometimes needed, because the usual method of choice, i.e., second harmonic generation measurements, is not always perfectly robust for such systems.

V. SUMMARY

In this paper we showed that $g$ factors of excitons in semiconductors (value and sign) can be determined by first-principles methods if the calculation of the orbital angular momentum $L$ is properly converged. For the considered two-dimensional materials hundreds of bands were required to obtain reasonable convergence, indicating that the basis set size is a critical numerical issue. For an individual Bloch state the calculation of $L$ suffers from the well-known band gap underestimation of density functional theory. However, the error in $L$ is approximately the same for electron and hole states and for excitons (which depend on the difference $\Delta L$) error cancellation enables quantitative calculations.

We applied the method to excitons in monolayers of semiconducting MX$_2$ (M = Mo, W; X = S, Se, Te) and interlayer excitons in MoSe$_2$/WSe$_2$ heterobilayers and obtain good agreement with available experimental data. The precision of our method allows to assign measured $g$ factors of optical peaks to specific transitions in the band structure and also to specific regions of the samples. This revealed the nature of various, previously measured interlayer exciton peaks. We further show that due to specific optical selection rules $g$ factors in van der Waals heterostructures are strongly stacking- and spin-dependent.

The presented numerical approach can be applied to a wide variety of semiconductors. Combined with $g$ factor measurements it might become a useful tool that helps to reveal the nature of optical excitations in semiconductors.

Note added. During the submission of this article three preprints on the calculation of exciton $g$ factors of TMD monolayers using first-principles methods appeared [68,79,81].

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