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An Account on BiVO₄ as Photocatalytic Active Matter

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CONSPECTUS: Photocatalytic materials are gaining popularity and research investment for developing light-driven micromotors. While most of the early work used highly stable TiO_2 as a material to construct micromotors, mostly in combination with noble metals, other semiconductors offer a wider range of properties, including independence from high-energy UV light. This review focuses on our work with $BiVO_4$ which has shown promise due to its small band gap and resulting ability to absorb blue light. Additionally, this salt's well-defined crystal structures lead to exploitable charge separation on different crystal facets, providing sufficient asymmetry to cause active propulsion. These properties have given rise to fascinating physical and chemical behaviors that show how rich and variable active matter can become. Here, we present the synthesis of different $BiVO_4$ microparticles and their material properties that make them excellent candidates as active micromotors. A critical factor in understanding inherently asymmetric micromotors is knowledge of their flow fields. However, due to their small size and the need to use even smaller tracer particles to avoid perturbing the flow field,



measuring flow fields at the microscale is a difficult task. We also present these first results, which allow us to demonstrate the correlation between chemical reactivity and the flow generated, leading to active motion. Due to the nontoxic nature of $BiVO_4$, these visible-light-responsive microswimmers have been used to study the first steps toward applications, even in sensitive areas such as food technology. Although these initial tests are far from being realized, we have to face the fact that a single microswimmer will not be able to perform macroscale tasks. Therefore, we present the reader with the first simple studies of collective motion, hoping for many new contributions to the field. The one-step synthesis of $BiVO_4$ clearly paves the way for studies requiring large numbers of particles. We predict that the combination of promising applications for a nontoxic material which is readily synthesized in large quantities will contribute pivotally to advance the field of active matter beyond the proof-of-concept stage.

1. INTRODUCTION

1.1. Light-Driven Micromotors

Visible light refers to a fraction of electromagnetic radiation that is perceived by the human eye, providing the raw data that, in turn, enables vision. On earth, sunlight is the main source of energy. Nature takes advantage of it through different mechanisms that transform the energy stored in electromagnetic radiation into chemical energy or mechanical (kinetic) processes. Because the energy content of light is easily variable by employing different wavelengths and intensities, light has proven to be an attractive energy resource to power-up and activate synthetic active colloids. Remote energy transmission and optical actuation have become a frequently used tool in many different approaches.¹ There are excellent reviews explaining the history of light driven active matter in-depth and under different perspectives,^{2,3} for which we will only summarize general information and focus on the contemporary developments on BiVO₄ here.

Since photoactivated colloids require light to excite chemical processes that generate the self-created chemical gradients, their motion can easily be activated by switching the light source on and off. One of the earliest photocatalytic microswimmer examples are Au@TiO₂ Janus particles.^{4,5} UV light illumination causes the generation of electron-hole pairs

in TiO₂, which are then separated by a Schottky junction, leading to electron transfer to the Au hemisphere. In short, the photoactivation of the photocatalytic material enables the catalytic decomposition of a fuel that forms a gradient which propels the swimmers, whereas in absence of irradiation, they simply perform Brownian motion.⁵ The control-enabling advantages of light over other energy sources have led to an increase in related research on microswimmer technologies. Soon, further materials besides TiO₂ and ZnO were developed that allow to switch to visible light. Possible pathways to visible-light-activated photocatalysts are, for example, the engineering of heterostructures or switching to semiconductors with a smaller bandgap.

1.2. Properties of Bismuth Vanadate

Vanadates have historically been used as a variety of pigments, or as test reagents for detecting illicit substances.⁶ As a mixed-

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Figure 1. Synthesis scheme and SEM images of polycrystalline $BiVO_4$ microparticles at pH 2–3, causing the formation of particles with spheroidal and rectangular and stacked morphologies. Adapted with permission from ref 16, 18. Copyright 2021 MDPI.

metal oxide, BiVO₄ has recently gained interest as catalyst for photoelectrochemical water splitting, as it exhibits favorable valence and conduction band potentials for water oxidation.⁷ Extensive research on the fabrication of BiVO₄ photoanodes for photoelectrochemical processes has been conducted over the past 20 years. In particular, the monoclinic crystalline structure is more appealing for photocatalytic processes, due to its short band gap (2.4 eV).⁸

The band gap type of BiVO₄ has been a topic of discussion: Zhao et al. found, in calculations, that the band gap between conduction band minimum and valence band maximum of BiVO₄ is indirect, but direct band gaps with slightly higher energies are also present.⁹ Walsh et al. confirmed the dominance of the direct band gap¹⁰ by good linear correlations between $(\alpha h\nu)^2$ and $h\nu$ in the Tauc plot of most experimental works.¹¹

The theoretically achievable solar to hydrogen (STH) efficiency of $BiVO_4$ in water splitting is promising for future applications.¹² However, the practically achieved STH remains at lower values to date, owing to the short carrier diffusion length and the slow hole transfer at the $BiVO_4$ /electrolyte interface.¹³

What makes $BiVO_4$ particles so relevant for microswimmer systems is the efficient spatial electron—hole separation upon illumination without the need of a cocatalyst.¹⁴ Monoclinic $BiVO_4$ crystallizes in an octahedral truncated bipyramidal shape, with mainly {010} and {110} facets being exposed. In 2013, Li et al. showed experimentally that photogenerated electrons are drawn to the {010} facet and holes to the {110} facet by photoreducing and -oxidizing a cocatalyst onto the different facets, and that the amount of oxygen evolution from water oxidation increased by a factor of 160 when particles with photodeposited Pt and Co_3O_4 were compared to plain $BiVO_4$ particles, compared to a factor of 9 for unselectiveley impregnated particles.¹⁵ In the field of light-driven micro/nanoswimmers, BiVO₄based systems are becoming ideal candidates to develop noblemetal-free micromachines without requiring complex fabrication techniques. Depending on their morphology, these devices have shown interesting swimming behaviors as a result of the photogenerated fluid flows, interactions with the substrate, and/or presence of external entities. In the following sections, we will describe our views on crucial points of motion mechanisms and interactions and give an outlook on future applications.

2. SYNTHESIS OF DEFINED BIVO₄ MICROPARTICLES

Syntheses of bismuth oxides are typically carried out in the acidic regime to solubilize bismuth ions, which otherwise react with water to form the insoluble salt bismuth oxynitrate (eq 1). This can be achieved using a variety of acids (eq 2), frequently HNO_3 .

$$Bi(NO_3)_3 + H_2O \rightarrow BiO(NO_3) + 2HNO_3$$
(1)

$$\operatorname{BiO}(\operatorname{NO}_3) + 2\operatorname{H}^+ \to \operatorname{Bi}^{3+} + \operatorname{H}_2\operatorname{O} + \operatorname{NO}_3^- \tag{2}$$

In the synthesis, the vanadate precursor NH_4VO_3 first forms vanadic acid (H_3VO_4), which eventually reacts with bismuth ions to form the insoluble mixed oxide BiVO₄. These particles can be either single-crystals or of polycrystalline nature, leading to different properties and advantages to either outcome. Within the hydrothermal synthesis approach, tuning the ratio of reagents, adapting pH values of the solutions,¹⁶ varying additives, or even the reaction medium, results in different morphologies.¹⁷

While other structures are available, we summarize typical syntheses that have been used to produce colloidal homogeneous $BiVO_4$ micromotors and their unique properties.



Figure 2. Left: Scheme showing the impact of addition of NaCl and SDS marked by arrows on crystal morphology and SEM images of BiVO₄ particles synthesized at different concentrations of NaCl and SDS: top: 0.05 M NaCl and 0.05 M SDS; second: 0.05 M NaCl and 0.005 M SDS; third: 0.05 M NaCl; bottom: 1.0 M NaCl. Right: XRD pattern and UV–vis absorption plot of the selected BiVO₄ sample. The monoclinic scheelite crystal structure can be clearly identified (Reference: PDF-Nr. 16-688, red). In the absorption spectrum, which was recorded with a dilute particle solution in water, it can be seen that light in the UV range up to the visible range is absorbed. Calculation of the band gap via the Tauc plot for a direct semiconductorgives a value of $E_g = 2.45$ eV. Reproduced with permission from ref 22. Copyright 2022 Wiley Online Library.

2.1. Polycrystalline Microparticles

In a typical synthesis of polycrystalline BiVO₄ colloids, the precursors $Bi(NO_3)_3$ and NH_4VO_3 are dissolved in a solvent mixture of ethanol, ethylene glycol (EG) and HNO₃, where EG executes two functions: it serves as a solvent that stabilizes BiO^+ ions and thereby increases the solubility of $Bi(NO_3)_3$ and acts as surfactant, stabilizing the {010} facet of the particles.¹⁹ At pH = 2, spheroidal particles were obtained, while if the pH was increased, the shape of the particles became rather rectangular, and at pH = 3, a cuboidal, almost star-shaped morphology is observed (Figure 1). In the presence of the chelating agent ethylenediaminetetraacetic acid more defined shapes, as well as single and double star-shaped particles are obtained, where the irregularly shaped, thin crystallites are stacked on top of each other, which leads to visible clefts between the layers.^{20,21}

The quite obvious change in particle shape over a rather narrow pH range from 2 to 3 can be explained by the concentration of H^+ ions, which influences crystal growth through the solubilization of bismuth ions.

Among the first reports using $BiVO_4$ as active matter, particles based on polycrystalline geometries were used that resemble stars or shuriken.^{20,21} These showed the peculiarity of

standing up on irradiation and swimming in an upright position.

2.2. Single-Crystals

Besides polycrystalline particles, single-crystals are also promising candidates for developing efficient light-driven microswimmers. This is due to a high electron—hole separation rate under illumination, resulting from the absence of grain boundaries. Shape and uniformity of single-crystalline BiVO₄ microparticles can be strongly improved by the addition of surfactants during the synthesis. An adapted protocol including the addition of NaCl and SDS (sodium dodecyl sulfate) as capping agents allowed the tuning of aspect ratios and yielded in rather homogeneous morphologies.²² Similar to the polycrystalline particles, XRD and DRS analysis revealed the monoclinic BiVO₄ structure and a band gap of 2.45 eV for the single crystals (Figure 2).

3. ASSESSMENT OF ELECTRON-HOLE SEPARATION OF THE MICROSWIMMERS BASED ON THEIR MORPHOLOGY

To investigate if this preferential crystal orientation along with the surface heterojunction commonly observed for $BiVO_4$ crystals between their {010} and {110} facets can serve as a



Figure 3. Scheme of charge separation on facets of $BiVO_4$ and identification by photodeposition. Reductive areas can be identified by reduction of noble metal ions like Ag^+ to Ag, while Co_3O_4 is preferentially deposited on oxidative areas. Reproduced with permission from ref 22. Copyright 2022 Wiley Online Library.



Figure 4. SEM images of (a, d) single-crystalline, (b, e) stacked, and (c, f) spheroidal BiVO₄ particles after photodeposition with (a-c) AgNO₃ and (d-f) CoCl₂ and NaIO₃. Reproduced with permission from refs 18, 22. Copyright 2022 Wiley Online Library.

means to separate excited charge carriers on the particle surface, photooxidation and reduction experiments were carried out with the different particle geometries. The principle of these experiments is shown in Figure 3.

To identify electron-rich facets using photodeposition, aqueous dispersions of $BiVO_4$ samples were illuminated with a halogen lamp while in the presence of Ag^+ ions, so that Ag^+ ions are reduced to $Ag^{(0)}$ by photogenerated electrons (eq 3).¹⁴ No hole scavenger was required, indicating that H₂O was oxidized to O_2 or •OH (eqs 4 and 5) on the side faces of the particles.

$$Ag^{+} + e^{-} \to Ag^{(0)} \tag{3}$$

$$2H_2O + 4h^+ \rightarrow O_2 + 4H^+$$
 (4)

$$H_2O + h^+ \to {}^{\bullet}OH + H^+ \tag{5}$$

SEM images of the selective photoreduction on different particle geometries are shown in Figure 4a–c. Given that the electrons preferably accumulate on the {010} facets at the top and bottom of the particles, these charges lead to localized deposition of noble materials. This behavior is consistent over the 3 different geometries but more pronounced for singlecrystalline BiVO₄ particles. For stacked particles deposition is mainly observed on the top and bottom areas, but also with less intensity into the clefts between the stacked layers. To confirm the presence of hole-rich facets, BiVO₄ particles have



Figure 5. (a) Orientation of single-crystalline, stacked, and spheroidal particles in a 0.1 wt % aqueous H_2O_2 solution in dark conditions and under irradiation with (385 nm) UV light. UV light irradiation causes a reorientation into an upright position for the single-crystalline and stacked particles, which was not observed for spheroidal particles. (b) Box plots of average instantaneous speeds of at least 25 single-crystalline (SC), stacked (PC stacked), and spheroidal (PC spheroidal) particles in a 0.1 wt % aqueous H_2O_2 solution under (385 nm) UV light irradiation. Summarized behaviors from refs 16, 18, 22, 23.

been illuminated in the presence of Co^{2+} ions, which can be oxidized to Co_3O_4 by photogenerated holes.¹⁵ As an electron scavenger, NaIO₃ was added to the solution. Under illumination, IO_3^- is reduced to I⁻.

$$IO^{3-} + 6e^{-} \to I^{-} + \frac{3}{2}O_2$$
 (6)

$$3\text{Co}^{2+} + 4\text{H}_2\text{O} + 2\text{h}^+ \to \text{Co}_3\text{O}_4 + 8\text{H}^+$$
 (7)

As observed before, the deposition on single-crystalline particles on the $\{110\}$ facets is much more pronounced compared to stacked particles. For spheroidal particles no photodeposited Co₃O₄ was detected (see Figure 4d-f).

To summarize, a clear orientation of the crystal structure is visible in single-crystalline microparticles, leading to preferentially exposed $\{010\}$ facets and charge carrier separation upon illumination.

4. CHARACTERIZATION OF MOTION

When aqueous dispersions of microparticles are mixed with H_2O_2 and illuminated with visible light, excited charge-carriers are created within the BiVO₄ microparticles which leads to respective reduction and oxidation processes of H_2O_2 . This duality results in the photogeneration of different ions and chemical species that diffuse at different rates, creating an asymmetrical chemical gradient around the particles that triggers their ballistic motion by phoretic mechanisms.

As the degree of asymmetry is mainly determined by charge carrier separation by the surface heterojunction between $\{010\}$ and $\{110\}$ facets in BiVO₄, different degrees of ballistic motion can also be expected for the three particle morphologies.

It is well-known that many Janus particle microswimmers swim in a fixed 90° angle to the substrate because the selfinduced flows around them lead to a reorientation.^{24,25} Similarly, we observe a reorientation for BiVO₄: in absence of suitable radiation, particles lie horizontally on their bottom face, while light irradiation causes a 90° rotation into an upright position, where the particles stand on one of their side faces (see Figure 5a).^{2,21,22} We believe that analogous to spherical swimmers, a combination of hydrodynamics, chemical fields, and second order effects are causing this behavior. For the spheroidal microparticles, observation of particle rotation around their long axis is nearly impossible in an optical microscope with two-dimensional resolution. Yet, all particles show increased activity and move in a certain direction, which is indicated by red arrows.

Comparing individual speeds, the single-crystalline BiVO₄ microparticles are fastest with a mean speed of 6.59 μ m s⁻¹, while polycrystalline particles are characterized by lower mean speeds (Figure 5b). Both, the stacked and spheroidal particles swim at speeds just over 4 μ m s⁻¹ in the ballistic regime, as verified looking at the slope of the bilogarithmic MSD plot for the time up to $\tau = 1$ s.

4.1. Flow Fields

Microscale motion is caused by interactions between the microswimmer surface, the solutes, and the substrate on which they are moving. These interactions cause fluid flows in the vicinity of the microswimmer, which eventually lead to the fluid executing forces on the microswimmer and the microswimmer back on the fluid, resulting in overall force- and torque-free motion.

Fluid motion on the microscale is described by the Navier– Stokes equations, with three basic solutions yielding possible flow fields. A stokeslet or force monopole is the simplest solution obtained for a particle driven by an external point force. A force dipole is the simplest solution of the Stokes equations that can describe active, self-induced motion and respects the boundary condition that the net force acting on an active particle is zero. Therein, two forces with the same magnitude point in opposite directions, and we can differentiate between two distinct cases: for outward pointing forces, the flow field is extensile and the resulting microswimmer is denominated pusher. In the opposite case, the forces point



Figure 6. (a) Schematic illustration of separated half reactions of H_2O_2 decomposition on the facets of single-crystalline BiVO₄ particles. H_2O_2 reduction, which consumes H^+ ions preferably takes place on the {010} facets, while protons are produced in the H_2O_2 oxidation on the {110} facets. (b) Light microscopy image of a typical flow field experiment. Reproduced with permission from refs 18, 22. Copyright 2022 Wiley Online Library.



Figure 7. Exemplary flow fields around attached (a) upright and (b) inclined microswimmers. Orientation and position of the swimmers are indicated by the yellow particle sketch. In the surrounding fluid of both colloids, flows from oxidative to reductive facets can be observed. Reproduced with permission from ref 22. Copyright 2022 Wiley Online Library.

inward, forming a contractile flow field and the swimmer is called a puller. 26

Experimentally, the flow field of microswimmers can be obtained by observing the trajectories of small passive tracer particles. Depending on the specific details, the measurements are referred to as particle image or tracking velocimetry, PIV or PTV, respectively, giving insights how the self-created solutes interact with the swimmer's surface, resulting in distinct flow profiles.²⁷ Due to the experimental difficulties on the microscale, the use of particle velocimetry on micromotors has been very limited and not yet extended beyond spherical microswimmers²⁸

For BiVO₄ microswimmers, these techniques were difficult to adapt, which was not only due to the deviation from spherical shapes for single-crystalline and stacked polycrystalline morphologies but also the origin of the activity is caused by photochemical reactions, leading to a completely different source distribution. To reliably track the flows, the choice of tracer particles is of the highest importance. Ideally, the tracer size should not be larger than 1/50 of the object diameter and the materials should not engage in chemical reactions. Au nanoparticles (Au NPs) proved to be a good material choice for BiVO₄ due to their high density (confining them mostly in one plane) and their excellent contrast in light microscopy. Despite their low mean diameter of (250 ± 22) nm, individual particles could be tracked in optical microscopy. Au NPs of that size also did not show significant catalytic activity for H₂O₂ decomposition.

The strong separation of oxidation and reduction sites in the single-crystalline $BiVO_4$ microparticles suggests that their motion pattern is dominated by self-electrophoresis, for more details see reference.¹⁸ Under illumination, H⁺ ions are produced at the {110} facets and consumed in a reduction reaction at the {010} facets (Figure 6a). Consequently, fluid flows from the side to the top and bottom faces of the microswimmer would be expected. In contrast to conventional self-electrophoretic microswimmers, the single-crystalline



Figure 8. (a) Zeta potential and Conductivity of stacked BiVO₄ particles at different pH values. Particles with positive zeta potentials perform horizontal motion (indicated by red area), while a reorientation and upright motion (indicated by blue area) was observed for negative zeta potentials. (b) Scheme showing immobilized HPDASCl on a glass substrate (left) and photoacid principle (right). Under UV illumination, an increased acid constant causes the dissociation of the molecule. Summarizing experiments from refs 18, 21. Copyright 2019 Wiley Online Library.

BiVO₄ colloids should therefore give rise to a source quadrupole in the focus plane due to their geometry: When in an upright position, two sides with $\{110\}$ facets are illuminated by light and act as proton sources, while the two $\{010\}$ facets consume protons (see Figure 6b).

Obtained flow fields are depicted in Figure 7a and b, comparing particles in an upright position (i.e., a 90° angle between the microswimmers and glass substrates) and an inclined position, revealing that tracer velocities and motion direction vary strongly. Still, common trends include the inward flows from the long side {110} toward the top and bottom {010} facets, from where the fluid appears to be pushed outward. The source quadrupole which is caused by electron driven reactions on the $\{010\}$ facets (along the *x*-axis) and hole-induced reactions on faces with {110} facets (along the y axis), is finally overlaid with hydrodynamic constraints and results in a flow field with similarities to a pusher-type force dipole. The asymmetry required for active motion is most likely ascribed to an overlap of intrinsic surface asymmetries and self-shadowing, which then translates into different amounts of created charge-carriers and subsequently into a more pronounced product gradient. These effects could be self-enhancing, which is underlined by the observation, that inclined microswimmers are often seen to move along rather straight trajectories.

Preliminary tests showed that flows caused by polycrystalline $BiVO_4$ microparticles were too small to be resolvable with the PTV methodology described.

4.2. Motion Mechanisms

Building on the charge carriers produced by photophysical interactions of irradiation with $BiVO_4$, it is reasonable to assume the motion mechanism in H_2O_2 to be electrophoretic. From a mechanistic and kinetic perspective, H_2O_2 degradation is a complex reaction: Under acidic and neutral conditions on

the surface of a suitable catalyst, hydrogen peroxide disproportionates into H_2O and O_2 (eq 8).

$$H_2O_2 \to H_2O + \frac{1}{2}O_2$$
 (8)

When charge carriers are available in the system, the disproportionation reaction can also be separated into an oxygen-forming oxidation half-reaction and a reducing half-reaction of H_2O_2 (eqs 9 and 10).

$$H_2O_2 + 2h^+ \to O_2 + 2H^+$$
 (9)

$$H_2O_2 + 2H^+ + 2e^- \to 2H_2O$$
 (10)

Looking at the equations above, it becomes apparent that two different species can create chemical gradients, leading to phoretic motion by different mechanisms: O_2 resulting in neutral self-diffusiophoresis, and H⁺, which also generates an electric field and therefore results in self-electrophoresis.^{29,30} While simple neutral diffusiophoretic models capture certain phenomena, many experimentally observed behaviors cannot be explained considering only neutral reaction products.³¹ Ionic diffusiophoresis considering also an endogeneous electric field, resulting from the diffusion of charged species, broadens the applicability.³² Electrophoretic micromotion was first described for bimetallic Au–Pt rods in dilute H_2O_2 .^{33,34}

The overall catalytic reaction is split into reduction and oxidation half reactions (see also eqs 9 and 10), happening at opposite sides of the swimmer body, with electrons migrating through it, causing a net displacement in a self-generated electric field, driven by the motion of ions and subsequent flows in a thin layer around the particle.^{29,35} In analogy to TiO₂ based microswimmers³⁰ we assume the dominant mechanism to be electrophoretic. This is underpined by the change in direction of particle motion upon an inversion of the zeta



Figure 9. (a) Scheme showing the spatially separated half reactions on the facets of $BiVO_4$ microcrystals. The oxidation of Dibenzylamine to *N*-Benzylidenebenzylamine is taking place on the {110} facets while O_2 is oxidized to H_2O on the {010} facets. (b) Scheme demonstrating the created gradients in H⁺ concentration and the resulting flows. (c) Box plots of average speeds of BiVO₄ microswimmers under two intensities of blue light (468 nm) and in different DBA concentrations (*x* axis is not to scale). Reproduced with permission from ref 37. Copyright 2022 Wiley Online Library.

potential as well as further environmental parameters such as the pH and conductivity.

4.3. Environmental Factors Affecting Motion

Conductivity. The conductivity of the solution is a crucial parameter to prove ionic or electrophoretic influences,^{31,36} since with increasing conductivity, the endogenous electric field around a microswimmer decreases according to Ohm's law. For self-electrophoresis, the swimmer speed is predicted to be proportional to this self-produced electric field,³⁵ while for self-diffusiophoresis the relations are much less clear, as factors like viscosity and catalytic activity are influenced by the ion concentration. A test for stacked BiVO₄ particles in increasing concentration of NaCl resulted in a steady decay of the swimming speed starting from a concentration $c_{\text{NaCl}} = 1 \times 10^{-6}$ M.¹⁸ Concentrations higher than $c_{\text{NaCl}} = 1 \times 10^{-2}$ M resulted in stuck colloids.

pH. The motion of $BiVO_4$ particles is strongly influenced by their surface charge, which depends on the pH value and the point-of-zero-charge. For stacked particles, we found that their motion mode changes from upright to horizontal depending on the pH values.^{2,21} Figure 8a displays the zeta potential and conductivity of stacked $BiVO_4$. From pH 5–9 the particles show a negative zeta potential, perform a reorientation and move in an upright orientation. However, if the pH is decreased and the zeta potential changes to positive values no reorientation and a horizontal motion is observed.

Based on this observation, we explored the possibility of locally influencing the pH in the vicinity of the particles. For that purpose, the photoacid 3,8-bis(N,N-dimethylsulfamoyl)-6hydroxypyrene-1-sulfonyl chloride (HPDASCl) was coupled to APTES and immobilized on glass substrates.¹⁶ Under UV light irradiation, the pK_a value of HPDASCl in water decreases from 5.7 to -2.4, which causes a decrease on local pH and possibly inverts the zeta potential of the BiVO₄ particles from negative to positive (see Figure 8b). Experiments were carried out in a dilute H_2O_2 solution at pH 6.2. Under these conditions, only upright motion was observed on unmodified glass substrates. However, on substrates with immobilized photoacid about 40%-80% of particles show upright motion and 12%-45% show horizontal motion. Additionally, we observed a third motion mode, that was called tumbling, where the particle either switches between horizontal and upright repeatedly or cannot be assigned to any of the two at all. On the other hand, the morphology of the microswimmers also affects their swimming behavior. A comparison between single and twin

star shaped $BiVO_4$ microswimmers showed that only the latter exhibited a stand-up motion upon light irradiation, attributed to a higher accumulation of the photogenerated products due to their concave morphology.

Chemical Fuels. The previous studies in dilute H_2O_2 under UV illumination showed a dependence on fuel concentration and the influence of reaction- and catalystdependent equilibria.¹⁸ These experiments ensured the comparability with existing studies of Janus particle swimmers and theoretical models. However, the highly oxidative character of the photogenerated species upon activation of photocatalytic microswimmers can also be used to target other types of reactions, e.g. the oxidation of dibenzylamine (DBA) to N-benzylidenebenzylamine.³⁷ Similar to the decomposition of H_2O_2 , the oxidation reaction produces H^+ while, at the same time, H⁺ ions are consumed in the reduction reaction (Figure 9a). Driven by the resulting flows, the particles moved in different concentrations of DBA in acetonitrile under visible light (Figure 9b and c), demonstrating that microswimmers can be propelled by an organic reaction in media different from those in aqueous solutions. Furthermore, we could also show, that nontraditional fuelling reactions, such as the photodeposition of metals from their respective salts, are suitable to propel BiVO₄ micromotors.³⁸ Noticeable therein is that because the gradient driving the motion is a negative one, the complex metal ions are removed from the solution and deposit on the BiVO₄ surface.

5. INTERACTIONS

5.1. Active-Active Interactions

The formation of larger groups, so-called schools or flocks, often provides living active matter with remarkable functions, such as protection or foraging benefits. Often, these interactions happen without external coordination or supervision, resulting in a profound lack of understanding of group dynamics. So-called "arising of spontaneous complex phenomena" has fascinated physicists for decades, but we are still far away from understanding. Here, the interaction among seemingly simple, artificial active matter holds promises to abstract and model some of these interactions, which brings them to the attention of behavioral biologists as well as scientists developing future applications.

Modular microswimmers (i.e., reconfigurable assemblies of multiple particles into one larger swimming object) are a specific subspecies where a defined geometry may open

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Figure 10. Light micrographs of assemblies of two $BiVO_4$ microparticles of the same morphology, as guides to the eye, trajectories over several time units are represented with red lines. Single-crystalline and stacked particles assemble along their side faces and circular trajectories are observed. Spheroidal particles assemble along their short axis and do not show increased circularity in their trajectories. Comparing particle behaviors from refs 18, 20, 21, 23. Copyright held by the authors.



(b) Experiment of large assemblies



Figure 11. (a) Experimental micrographs (upper line) and simulation snapshots (lower line) of modular swimmers with recurring geometries consisting of two to four microparticles. Insets in experimental images show exemplary SEM images of the respective geometry for visualization. Inset scale bar is 2 μ m. (b) Collective assemblies of star shaped BiVO₄ micromotors induced by switching the light on/off. Scale bars: 10 um. Reproduced with permission from refs 20, 23. Copyright 2019 and 2020, respectively, American Chemical Society.

possibilities to incorporate different functionalities into one swimmer. Generally, these assemblies are to be understood as an intermediate between single microparticles and swarms thereof.^{39,40} Previously, these mostly spherical building blocks were made of a single material without inherent structural anisotropy and therefore immotile when isolated, but upon assembly asymmetry is increased and motion is enabled.^{39,40} For BiVO₄ microparticles, their inherently anisotropic semiconductor nature enables motility also in individual particles, but their nondipolar source geometry seems to favor attractive interactions. We observed attraction among all BiVO₄ microparticle geometries: single-crystalline and stacked microparticles attract each other mainly along their side faces (Figure 10), which under illumination are hole-rich and seem to create mainly inward flows. As the particles are not permanently bound to each other, the assemblies reconfigure and break apart repeatedly under illumination. If the exciting UV light is turned off, all light-induced forces immediately vanish, which causes the particles to disassemble.

For spheroidal microswimmers, an increased tendency to form larger assemblies, preferably along their side faces, was observed. Upon contact with other spheroidal microparticles, these assemblies grow into motile, recurring geometries with up to four microparticles, that move actively (Figure 11a).

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Figure 12. (a) Intrinsic interactions of $BiVO_4$ microswimmers with yeast cells under visible light irradiation, involving a swimming trajectory change to capture and transport a yeast cell. (b) Schematic illustration of yeast cell removal by $BiVO_4/Fe_3O_4$ microswimmers from an unfiltered beer under combined light/magnetic fields. It also shows an optical microscope image of a beer sample before and after treatment. Reproduced with permission from refs 20, 46. Copyright 2019 American Chemical Society and 2020 Chemistry Europe, respectively.

Similarly, star shaped $BiVO_4$ particles reversibly form larger clusters (Figure 11b).

For spheroidal microswimmers, the interactions were found to happen on the long side of the ellipsoids, i.e., the moving direction. To theoretically reproduce this behavior, interactions were implemented in solid particles, constructed from partially overlapping spheres with catalytic activity implemented along one side of the spheroids. In consequence, a chemical concentration field $c(\mathbf{r}, t)$ is produced at a rate k_0 . All colloids in the system respond to this field, which effectively causes self-propulsion and attraction between colloids.

This model produces assemblies consisting of three and four microparticles in two different configurations: linear "caterpillars" and a more compact configuration, and reproduces the trends in speed. called "triangle" and "rectangle". The speed of assemblies decreases with increasing amount of particles in experiments as well as in simulations, which is likely related to the change in the chemical gradient around larger assemblies in relation to their drag force. Toward a complete understanding of collective interactions of inherently anisotropic micromotors, including the larger, frequently immotile assemblies that were observed at long illumination times, further experiments are required.

5.2. From Active–Passive Interactions to Synergies with Microorganisms

Considering active matter in Nature, collective interactions are fascinating and highly relevant for certain systems such as bacteria within biofilms, while for other swimmers, such as sperm or algae, the probability to encounter obstacles or alien species is much higher.

Electrostatic and phoretic interactions are some of the most common causes inducing cargo transportation by photocatalytic microswimmers.⁴¹ Star-shaped BiVO₄ micromotors have demonstrated a high driving force enabling the active particles to transport objects of different sizes, including 2 μ m polystyrene (PS) particles, but also bigger objects that exceed the size of the micromotors. This type of micromotor can either rotate in place or show directional motion. It has been reported that, by rotating, they can easily grab/release micron-sized polystyrene particles by switching the light ON/OFF. The capabilities of BiVO₄ micromotors to interact with living microorganisms were also investigated, but specific behaviors are discussed within the next section.

6. APPLICATIONS

Photocatalytic systems can be used in a plethora of applications ranging from selective oxidation to reductive reactions. Ideally, smart micro/nanoswimmers should perform specific tasks in a programmable and autonomous way, i.e., without requiring constant human assistance for motion control. In Nature, similar mechanisms are known for living microorganisms, which are mediated by chemical signaling according to the environment conditions. Due to the ability to outswim diffusion, environmental remediation is one of the top applications for micromotors, but the scaling discrepancies between tiny colloids and huge problems are dramatic. Using larger collectives of microswimmers can be a way to solve this discrepancy. Self-propelled photocatalytic micromotors have been used mostly for degradation of organic pollutants and recently, for removing plastic-based pollutants.⁴²⁻⁴⁴ This oxidation reaction is mainly mediated by the generation of highly reactive species such as hydroxyl and superoxide radicals. In this section, we will discuss the main proof-ofconcept applications of BiVO₄ microswimmers that have been reported so far and offer a perspective on what might be required to push them beyond 'proof-of-concept'.

6.1. Degradation of Organic Pollutants

Similar to the majority of light-driven photocatalytic micromotors, $BiVO_4$ has the ability to oxidize organic pollutants upon illumination. The coupling of $BiVO_4$ with a second photoactive material is beneficial to enhance electron-hole pair separation, leading to higher photocatalytic activity than bare $BiVO_4$. Dong et al. have reported a novel approach based on cooperative interactions between two types of micromotors, involving BiVO₄ and BiVO₄/graphene oxide micromotors with spherical and ellipsoidal morphology, respectively. The assembly/disassembly between micromotors can be easily controlled by adjusting the light intensity, due to an interplay between self-diffusiophoresis and electroosmotic mechanisms. Such cooperative interactions were used for the removal of rhodamine B as a model pollutant. The mixture of BiVO₄ and BiVO₄/graphene oxide micromotors achieved the highest pollutant removal efficiency in comparison with only BiVO₄ micromotors. This has been attributed to the aggregation/ disassembly among both types of motors under different light intensities that lead to a higher stirring of the liquid media, enhancing the mass transfer among RhB and the micromotors.

6.2. Antimicrobial Activity

Besides chemical contamination in wastewater, the presence of pathogenic microorganisms is one of the major causes of waterborne infections associated with drinking-water safety. Photocatalytic-based microswimmers can act as pathogenneutralizing devices, due to the in situ photogeneration of reactive oxygen species. Photocatalytic BiVO4 star-shaped swimmers are able to autonomously swim toward yeast cells (funghi), capture them by phoretic interactions and transport them under constant light irradiation.²⁰ The surface of BiVO₄ was not functionalized nor the swimming direction was manipulated by magnetic fields, indicating their autonomous behavior and intrinsic capabilities for capturing microorganisms. The BiVO₄ microswimmers showed the same capability of capturing yeast cells in the presence of other type of microorganisms, such as E. coli. Although some bacteria could attach to the BiVO4 motors, it did not hamper their ability to swim toward yeast cells and attach to their surface. Even though this behavior has been ascribed to a combination of several factors, including roughness surface, hydrophobicity, and surface interactions, the exact mechanisms of such interactions are not well understood. Before any real applications can be tackled, this lack of understanding needs to be addressed.

6.3. Food Industry

Yeast plays a key role in the manufacturing of food and alcoholic beverages. Therefore, the spontaneous interactions between artificial microswimmers and yeast cells can have a plethora of applications. Common approaches for yeast removal after the fermentation processes rely on pasteurization or filtration treatments. BiVO₄ star-shaped microswimmers modified with magnetic nanoparticles were shown to remove yeast contamination in real samples, such as alcoholic beverages (see Figure 12) because the self-propelled BiVO₄ microswimmers autonomously bind to yeast cells. Owing to their dual optical/magnetic response, the resulting assembly micromotor/yeast can be easily removed after treatment, simplifying the whole removal process. The treatment of unfiltered beer with BiVO₄/Fe₃O₄ micromotors resulted in almost 100% yeast removal. More importantly, there were no differences in the organoleptic properties of beer before and after treatment. The leakage of Fe and V ions was also examined, finding only remaining traces that are allowed in drinking water. Therefore, this work represents a simple approach to overcoming yeast spoilage in alcoholic beverages for applications in microbreweries that can be extended to other food-related processes.

There are diverse applications for BiVO₄ micro/nanomotors that are yet to be explored. For instance, BiVO₄ is a wellknown yellow pigment with antifouling properties.⁴⁷ Moreover, it has been widely explored for photoelectrochemical water oxidation, mainly decorated with cocatalysts or a second semiconductor. On the other hand, owing to the mild oxidizing power in comparison to TiO_2 , it is also a very interesting material for the generation of liquid solar fuels from methane, such as methanol.⁴⁸ Therefore, the improvement of the micromixing by the active self-propulsion of BiVO₄ micromotors might result in superior efficiencies of such challenging selective oxidations or add new dimensions for previous wellestablished reactions. Another interesting approach involves their coupling with enzymatic components for potentially driving photobiocatalytic reactions or enhancing and controlling the motion and functionalities of enzyme-based micro/ nanoswimmers.49,50

7. CONCLUSIONS AND OUTLOOK

In this Account, we discuss the importance and versatility of $BiVO_4$ not only as a material for photoelectrochemical processes but also as an efficient photocatalyst for developing efficient light-driven microswimmers. Owing to its comparatively small bandgap, $BiVO_4$ microswimmers can propel under visible light irradiation, which is desirable for a variety of applications that otherwise are hampered by the use of harmful UV wavelengths. Therefore, this feature enables the application of $BiVO_4$ microswimmers in water disinfection under biocompatible light wavelengths.

Besides, BiVO₄ can be synthesized in multiple shapes by easily scalable techniques, mainly involving co-precipitation and hydrothermal treatments. Depending on the morphology, from spherical, star-shaped, to single-crystals, BiVO₄ microswimmers show interesting swimming behaviors. For instance star-shaped micromotors show an unique stand-up motion behavior, due to chemical gradient build-up between the particle and the substrate. On the other hand, single-crystalbased micromotors show enhanced speeds in comparison with polycrystalline ones, which is attributed to an spatial electronhole separation of the photogenerated excitons in each crystal facet. Therefore, unlike common Janus microswimmers based on metal-semiconductor heterojunctions, BiVO₄ based devices can effectively separate electron-hole pairs without requiring the inclusion of an additional component in the micromotor structure. In this way, we can skip the use of large and expensive equipment, such as metal evaporators.

Their motion is affected by the surrounding pH, properties of the substrate, and type of fuel, offering more options for their actuation and manipulation. BiVO₄ microswimmers have also the ability to interact with each other and with the surroundings, which has led to emerging collective interactions, such as clustering-like behaviors and/or self-assembly, which are common for polycrystalline spheroidal particles. Such interactions have been useful to develop different microswimmer-based configurations and perform cargo transportation of passive particles (e.g., polystyrene) and living microorganisms. When yeast cells were selected as model microorganisms, the polycrystalline stacked BiVO₄ swimmers moved toward the yeast and captured it, involving even change of their swimming direction. However, further studies including flow field analysis are required to fully understand such interactions, which will enable the development of more sophisticated and controllable photoactive microdevices for

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biomedical applications such as photodynamic therapy. The low price and toxicity associated with this material raise expectations for applications beyond proof-of-concept for this convenient material. However, more systematic studies dealing with the reusability of these micromotors in multiple tests and an investigation of possible leaching of metals upon irradiation should be considered in the future. Moreover, the decoration of BiVO₄ with other semiconductors and/or materials would be beneficial to induce their motion in H_2O_2 -free environments to avoid adding extra pollution in the liquid media.

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/accountsmr.3c00021.

Movie showing single crystalline, polycrystalline stacked, and polycrystalline spheroidal particles (AVI)

Movie showing yeast capture by a particle (MP4)

Synthesis and characterization of BiVO₄ particles (PDF)

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Notes

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Katherine Villa received her PhD degree in Chemistry from the Autonomous University of Barcelona, Spain. After two research positions at the Catalonia Energy Research Institute and the Institute for Bioengineering of Catalonia, she joined the Center for Advanced Functional Nanorobots at the University of Chemistry and Technology, Prague (Czech Republic), where she worked as a senior scientist for 3 years. Currently, she is leading a research group on light-driven nanomotors and solar fuels at the Institute of Chemical Research of Catalonia (ICIQ). Her research interests include photocatalysis, micro/nanomotors, nanomaterials, renewable energy, and environmental remediation.

Juliane Simmchen studied chemistry in Dresden and Naples and obtained her PhD in nanotechnology at the Autonomous University of Barcelona. She then spent a short postdoc in Samuel Sanchez's group at the MPI for Intelligent Systems. On her return to Dresden, she was awarded a prestigious Freigeist fellowship to set up an independent research group in physical chemistry. In 2022 she moved to Strathclyde as a Reader in Materials Chemistry.

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