BODIPY- and Porphyrin-Based Sensors for Recognition of Amino Acids and Their Derivatives

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Abstract: Molecular recognition is a specific non-covalent and frequently reversible interaction between two or more systems based on synthetically predefined character of the receptor. This phenomenon has been extensively studied over past few decades, being of particular interest to researchers due to its widespread occurrence in biological systems. In fact, a straightforward inspiration by biological systems present in living matter and based on, e.g., hydrogen bonding is easily noticeable in construction of molecular probes. A separate aspect also incorporated into the molecular recognition relies on the direct interaction between host and guest with a covalent bonding. To date, various artificial systems exhibiting molecular recognition and based on both types of interactions have been reported. Owing to their rich optoelectronic properties, chromophores constitute a broad and powerful class of receptors for a diverse range of substrates. This review focuses on BODIPY and porphyrin chromophores as probes for molecular recognition and chiral discrimination of amino acids and their derivatives.

Keywords: molecular recognition; covalent recognition; chiral discrimination; BODIPY; porphyrin tweezers

1. Introduction

For several reasons, the visible light became an important aspect of research focusing on specific interactions with the matter, which can find a wide scope of applications. The inexhaustible access to visible light opens many paths for searching for cost-efficient light-activated catalysts or systems properly interacting with specific wavelength (λ), essential for optoelectronic applications. The generality of the light also drives the activity towards formation of systems that can involve, e.g., intramolecular transformation leading to substantial change in absorbance or emission indicating a presence of specific factor. The analytical tools based on application of the visible light are constantly explored because of straightforward detection of the change even with the eye. The construction of skeleton responding to specific factors requires precise planning and designing of structural motifs that would react with and eventually modify the observed outcome, starting to behave as a molecular probe. Covalent recognition is the ability of designed molecular probes to react towards target skeletons with the formation of a covalent bond. As a result, a different behaviour, especially the optical response, can be used to monitor the presence of the target molecule in the medium that is tested. Molecular recognition is a specific tendency of compounds to selectively bind to each other through non-covalent interactions [1–3], such as metal coordination [4], hydrogen bonding [5], hydrophobic interactions [6], van der Waals interactions [7], π–π stacking [8] or electrostatic interactions [9]. The term “chiral recognition” refers to an event of molecular recognition, in which a receptor exhibits higher affinity towards one of existing enantiomers of guest molecule. If the host molecule is chiral, the formed
host-guest complexes are diastereoisomeric and therefore possess sets of different properties, enabling their easy separation, as well as differentiation in $^1$H NMR spectroscopy (chiral discrimination).

In the past decades the BODIPY chromophores (Scheme 1, top) as well as porphyrins (Scheme 1, bottom) or porphyrinoids have found widespread applications as sensors for detection of a broad scope of substrates based on several straightforward detection channels, including UV/Vis, fluorescence and circular dichroism (CD) spectroscopies. These techniques provide a quick and accurate response in addition to requiring minimal amounts of analyte. It usually needs a properly defined optic response followed by high extinction coefficients and effective fluorescence with high quantum yield. The linear construction of BODIPY leads to sharp absorption and highly efficient emission observed, what opens a potential for application of those skeletons as optical probes. On the other hand, owing to the unique properties of porphyrinoids, they are interesting candidates for molecular [10] and chiral [11] recognition with multi-channel and straightforward UV/Vis, fluorescence and CD detection [12,13]. The macrocyclic character of porphyrin (Scheme 1, bottom) shows another aspect of oligopyrroles. Strongly conjugated $\pi$-clouds with a diatropic circuit operating within the porphyrins $18\pi$ electron path can be also applied in NMR detection, eventually showing a significant difference in registered chemical shifts. On the other hand, their rich optical properties, such as intense absorption in the Soret region (400–430 nm) and fluorescence, gives further possibilities for using them as probes as their strong and narrow absorption and fluorescence bands allow detecting even a slight perturbation caused by a guest binding.

![Scheme 1](image)

**Scheme 1.** Representation of the difference in the target recognition between: BODIPY probe (1a); and [18]Tetraphyrin(1.1.1.1) probe (1b) (porphyrin). The yellow triangle represents the target molecule, the light blue triangle represents a leaving group.

The abundance of molecular recognition phenomena in living organisms [14] is one of the many reasons this field of interest draws attention of many researchers. Herein, we present recent reports on BODIPY- and porphyrin-based sensors for the recognition of amino acids and discuss their respective applications. The importance behind the recognition of amino acids lies on the fact that those molecules are essential in the human body for many physiological processes such as protein synthesis, detoxification, metal binding and many more [15]. The over- or under-expression of those natural derivatives is usually strictly related to different pathologies, hence the ability to detect such abnormalities is very useful especially for the screening of diseases in patients. Moreover, recognition of amino acids and other natural derivatives is a powerful tool that, if combined to the peculiar optical properties of BODIPY molecules [16,17], finds applications in bioimaging which is a non-invasive diagnostic tool of extreme importance [18].

The detection of other natural derivatives based on amino acids, such as G-proteins [19], neurofibrillary tangles [20] and bio-thiols [21], are strongly related with many cellular functions and pathologies and are also mentioned in this work.
2. BODIPY Probes

BODIPY (BOron(III) DIPYrromethene) is a class of chromophore formed from two pyrrolic units linked by a methylene bridge (Schemes 1 and 2) in addition to coordinating electron-deficient boron(III). The planarity of those molecules jointly with the presence boron(III) give structures with extraordinary optical properties that are fascinating the research world for many decades. Extensive investigation and studies on the synthetic aspects of those derivatives have been carried out and allowed the design of many chromophores with versatile electronic properties. Those skeletons have found applications as probes since they have a peculiar optical behaviour presenting very sharp emission bands with a quantum yield close to a unit and more importantly even small changes in the molecular structure can trigger drastic shifts in the fluorescence spectrum [22–24]. Because of this, BODIPY probes are widely used as molecular recognisers for the target with covalent interaction; however, the involvement of non-covalent interaction was also reported as an efficient process allowing efficient recognition. The most important features of those molecules rely on the fact that they present narrow absorption and emission bands together with high quantum yields approaching a quantitative fluorescence [25] forming skeletons potentially applicable in several fields including the recognition of specific targets. The BODIPY core was extensively explored with an involvement of specific positions (Scheme 2), eventually leading to extraordinary behaviour including tuning of the optical properties with a modulation of absorption and emission wavelength in both directions–bathochromically [26] and hypsochromically [22]. Most of the modifications showing the skeletal dependence of the optical properties are based on covalent interaction but the probing behaviour needs a response to specific initiator or target potentially available with the applied structure.

![Scheme 2. Examples of possible modification on the BODIPY core.](image)

One of the most fascinating ability for those chromophores is the change in the optical response triggered by simple initiators such us pH, presence of anions and other factors [27]. Because of this behaviour, BODIPY molecules are very useful optical tools, finding many applications including the probing of, e.g., metals [28] or nitrate [29]. One other field where BODIPY derivatives are involved is the bioimaging and detection of specific targets found in the human body for many different reasons and applications [18].

The detection of those components can be achieved in different ways but, in general, a shift or the ON/OFF switch of absorption/emission is the final outcome in the recognition of a target.

A recent work published by Farinone et al. in 2020 showing the optical behaviour for BODIPY derivatives connected to different amino acids is the example to introduce the topic of BODIPY-amino acid derivatives [23]. The hybrid derivatives were obtained through nucleophilic aromatic substitution in the meso-position of the BODIPY chromophore, as shown in Scheme 3.
Those observations show how the optical properties of BODIPY skeleton can be modulated by a simple emission and absorption of the probe (Scheme 4). The sensing of the target molecules can be achieved towards the formation of a direct bond between the target and the BODIPY sensor with involvement of different positions of the fluorophore core and usually follows the chemistry of a nucleophilic aromatic substitution. It usually requires a precise designing of the mother skeleton and specific location of good leaving group either at \( \alpha \)-or \( \alpha' \)-position. After the substitution (Schemes 3 and 4), a novel BODIPY-target hybrid structure is obtained with optical properties strongly different from the starting materials making the detection of the target molecule easy and reliable. This type of interaction between the probe and the target can trigger different type of responses, which were divided into two main categories: the activation in the emission of the probe (or, in some cases, the deactivation in the probe’s fluorescence) and the shift in emission and absorption of the probe (Scheme 4).

### 2.1. Covalent Interaction (Recognition)

Based on the type of heteroatom involved in the \( \text{meso} \) bond (3x), hence what type of amino acid was used, a difference in the electronic properties was recorded with shifts in both emission (range, \( \lambda_{\text{max}} = 452–535 \text{ nm} \)) and absorption (range, \( \lambda_{\text{max}} = 407–519 \text{ nm} \)). The most red-shifted for both absorption and emission was observed for 3f and the biggest hipsochromic shift was recorded for 3d. Those observations show how the optical properties of BODIPY skeleton can be modulated by a simple modification making this chromophore a versatile example of the optical probe.

**Scheme 3.** The amino acids used in this work are alanine, glycine, lysine, \( p \)-amino-phenylalanine, tyrosine and cysteine.

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**Scheme 4.** Representation of the two approaches in the recognition through covalent interaction observed for BODIPY chromophore. \( R_1 \), group with fluorescence quenching properties; LG, leaving group such as \(-\text{Cl} \) or \(-\text{SMe} \).
2.1.1. Emission Triggering

Depending on the planned recognition, specific response modulating optic properties are required. The potential application to biological tissues very often requires a long emission wavelength available for specifically modified BODIPY skeletons. In 2011, Shao et al. reported on a BODIPY-based molecular probe with specific detection of cysteine among the biological thiols [30]. The structure of the chromophore was designed as a sensor 4 (see Scheme 5) with a 2,4-dinitrobenzenesulfonyl (DNBS) protecting group connected to the phenol ring present on the α position of the BODIPY through an ethylene bridge giving a mute in emission skeleton 4.

\[ \text{Scheme 5. Preparation of the BODIPY probe specific for cysteine and recognition.} \]

The recognition of the target molecule results in the unmasking of the structure 4 and eventually an appearance of intense fluorescence in the red region at 590 nm [30]. The derivative 5 presents a red shifted emission because of the specific extension of delocalisation along the α position.

Another example based on the enhancement of the emission toward the elimination of a protecting group with quenching properties was reported by Wang et al. in 2018 [31]. They designed a BODIPY probe (6) (Scheme 6) for the specific recognition of glutathione (GSH). The sensor was functionalised with a 2,4-dinitrobenzenesulphonyl group as a fluorescent quencher but more importantly as a good leaving group reactive toward thiols. As expected, 6 presents very low emission (with a quantum yield of $\Phi = 0.03$) at 656 nm. The probing behaviour of 6 was tested in a THF/PBS 1:1 solution by reaction with various natural amino acids: lysine, asparagine, histidine, phenylalanine, tryptophan, glutamic acid, asparagine and cysteine, as well as homocysteine and glutathione. Cys and Hcy enhanced the fluorescence response with a quantum yield reaching 24% and 12%, respectively, at 659 and 657 nm while with GSH it reached 48% at 656 nm. Considering the differences in the intracellular concentrations of those natural derivatives, the response of the probe is remarkably selective toward GSH when tested on MCF-7 cells. All other amino acids did not affect the efficiency of the fluorescence emission.

The mechanism of recognition of GSH by the BODIPY probe is shown in Scheme 6 displaying the deprotection of the 2,4-dinitrobenzenesulphonyl moiety.

A covalent interaction between the BODIPY probe and the target molecules resulting in the activation of the chromophore emission was also presented in 2017 by Wang et al. [32]. In this work, they prepared a probe that can distinguish cysteine and homocysteine over GSH. They designed and synthesised a probe (8) featuring p-aminophenylthio and cyano groups attached to the BODIPY moiety. The decision on the presence of those groups was inspired by previous documented works that pointed out how specifically a p-aminophenylthio moiety can detect natural thiols.
The presence of sulphur from cysteine attached to the meso-position of BODIPY skeleton was reported as bathochromically shifting factor [23], but, after treatment with Cys (10), the absorption peak shifted hypsochromically by ~60 nm to 480 nm with a dramatic change in fluorescence efficiency eventually observed at 549 nm. The hypsochromic shift of BODIPY absorption was reported for meso-nitrogen substituted skeletons of chromophores [33] further shifted by deprotonation of meso-NH group [22]. A similar behaviour was detected in the case of Hcy with the formation of 9 (Scheme 7).

In contrast, upon treatment with GSH, only a slight enhancement in the fluorescence was detected with an increase in intensity of 18% when compared to Cys and Hcy, indicating that the probe is successfully discriminating Cys and Hcy over GSH. The proposed mechanism of interaction of the BODIPY with cysteine and homocysteine (Scheme 7) incorporates formation of 5- and 6-membered rings obtained upon nucleophilic attack of the thiol groups. In the case of glutathione, the formation of such derivatives is possible but not favoured since the molecule is much bulkier when compared to the two others.

One recent example of BODIPY-based amino acid sensor was presented in 2020 by Jeon et al. [34]. They designed and developed a BODIPY sensor with high affinity for amino groups by introducing at the meso-position a good leaving group with an active esters type of bond well known from peptide synthesis (Scheme 8).
was eventually used in a bioimaging application to sense lysine in PC3 cells showing good and
with a drastic decrease of fluorescence [35].

They synthetised a BODIPY-based probe (12) and the other one with a hydroxysuccinimidyl substituent (13). They both present very high recognition ability toward volatile amines such as ammonia, methylamine and dimethyl amine. While 12 was demonstrated to mostly find application in detecting amines in organic solutions and their vapours in solid states, 13 was investigated in water solutions toward the recognition of biological amines such as lysine residues on proteins, antibodies and nucleic acids. The recognition tests were carried out in PBS buffer (10 mM, pH = 7.4) containing 1% acetonitrile at room temperature. The probe presented an absorption peak at 622 nm, while the emission, recorded at 648 nm, was very weak (Φ = 0.001) because of aggregation-caused quenching. Upon stepwise addition of L-lysine, the absorption band rises to 517 nm with a blue shift of 105 nm because of the less electron withdrawing character of the \( \epsilon \)-amino group and the \( \epsilon \)-nitrogen substituted skeletons of chromophores [33] further shifted by

\[ \text{Lysine addition: } \text{Absorption peak at 622 nm, emission at 648 nm, } \Phi = 0.001 \]

The recognition of the amino acid (lysine) shows a PET effect.

A similar recognition behaviour was presented in 2016 by Das and co-workers. In their work, the aza-BODIPY probe 15 (Scheme 9) reported by Xiao and co-workers selectively responds to lysine with a drastic decrease of fluorescence [35].

The aza-BODIPY probe 15 presents an absorption maximum at 732 nm and a strong emission in the NIR region at 753 nm in dichloromethane. After incubating the probe with lysine in a 1:4 DMSO/water mixture (pH = 7.3), the fluorescence intensity strongly decreased because of the photo-induced electron transfer (PET) shown in Scheme 9 from structure 16. The selectivity for the target was demonstrated testing different amino acids and bio-thiols, which, after incubation with the probe, did not show a change in the fluorescence intensity. Because of this specific behaviour, the BODIPY-based probe was eventually used in a bioimaging application to sense lysine in PC3 cells showing good and promising results.

A similar recognition behaviour was presented in 2016 by Das and co-workers. In their work, they synthetised a BODIPY-based probe (17, Scheme 10) which exhibits specific recognition abilities for lysine over other amino acids [36].
The specificity towards recognition of cysteine over homocysteine and glutathione is presented in Scheme 12. The structural similarities (nucleophilic thiol and amino group) in the architecture of Hcy and GSH with Cys should not preclude a cascade reaction eventually leading to an optical
response. However, the limiting step for Hcy and GSH is the formation of the cyclic transition state, which will generate six- and ten-membered rings, respectively. This kinetically unfavoured step makes the cascade reaction very slow for Hcy and impossible for GSH, allowing a high specificity for cysteine.

Scheme 12. Specificity of the recognition toward cysteine is explained according to structural differences.

In 2019, Avellanal-Zaballa et al. presented a BODIPY probe designed with functionalities responsive to thiol containing amino acids [38]. The sensor presents two $\alpha,\beta$-unsaturated esters on both $\alpha$ positions of the BODIPY core (22, see Figure 1) as the reactive points that are able to react with the nucleophilic groups of the sensed targets. The formation of the probe was possible starting from a di-formylated derivative sequentially converted to the unsaturated ester through Wittig reaction.

Figure 1. Structure of the BODIPY sensor 22 specific for detection of cysteine, homocysteine and glutathione.

Because of the extension of delocalisation through the $\alpha$ positions of the sensor, an absorption band is present at 590 nm and an emission band is recorded at 600 nm. After treatment with any of cysteine, homocysteine or glutathione, the absorption band is gradually replaced by two blue shifted peaks at 550 and 515 nm; the emission is similarly shifted to shorter wavelengths at 560 and 525 nm. The tests were carried out in a 1:1 mixture of ethanol/HEPES to mimic the physiological media. Even at low concentrations, the detection of targets (micromolar) was possible by monitoring the fluorescence; hence, the detection of bio-thiols is easily accessible using the described probe.

Intermolecular Displacement

In some specific cases where a shift in absorption and emission is detected upon covalent interaction between the probe and the target, the mechanism for the change in the electronic properties is based on an intermolecular displacement.

In 2018, Zhao and co-workers reported a BODIPY chromophore applicable for specific recognition for cysteine in human plasma [39]. In this case, it presented a strong emission and with no protecting groups in the architecture. The detecting approach starts from formation of cysteine derivative 24 (Scheme 13) that undergoes intermolecular transformation changing the optical behaviour with the formation of 25.
The probe 23 was chosen because of its stability in buffer solutions and the presence of a good leaving group at the meso position. Moreover, the structure was designed as an asymmetric molecule with steric substituents favouring a ratiometric response. The fluorescence spectrum was recorded in a CH$_3$CN/PBS buffer (pH = 7.4) at 25 ºC displaying main absorption and emission at 491 and 515 nm for probe 23. After addition of cysteine, a nucleophilic aromatic substitution took place in the meso-position and a ratiometric response was observed with a dramatic shift both on absorption and emission with maxima recorded at 412 and 470 nm. The strong shift in the emission is not explainable with the presence of a sulphur at the meso position; according to other reports presenting dissimilar properties [23,33]. Indeed, due to the intramolecular displacement described in Scheme 13, the amino group replaced the sulphur function giving and explanation to the strong shift.

The specificity of this probe relies on the fact that upon treatment with another thiol rich natural derivative, glutathione (GSH), the absorption peak at 491 nm rapidly decreased and a red-shifted peak at $\lambda_{\text{em}} = 509$ nm was formed. In addition, a new emission wavelength was measured at 537 nm and both optical shifts were not ratiometric and support the formation of a thioether intermediary, 26; indeed, the intramolecular displacement is not available for GSH, as shown on Scheme 13.

Another example was presented in 2012 by Niu et al. [40]. The focus of their work was the ability to selectively detect the presence of glutathione (GSH) over cysteine and homocysteine. To achieve this, they designed a BODIPY-based probe (27, Scheme 14) substituted with a chlorine atom on the $\alpha$ position as a good leaving group and a triazole on the other $\alpha$ position to enhance the reactivity toward nucleophilic aromatic substitution because of the strong electron withdrawing character of this substituent. Moreover, the BODIPY derivative presents good solubility in water and a defined emission peak at 556 nm. The fluorescence response of the probe was tested in an aqueous HEPES buffer (20 mM, pH = 7.4) containing 5% acetonitrile at 37 ºC with stepwise addition of GSH. As a result, the $\lambda_{\text{em}} = 556$ nm emission peak was gradually replaced by a new band at 588 nm with a fluorescence ratio proportional to the concentration of GSH and to the formation of derivative 28.
With Cys, after the initial addition of the nucleophile, a new emission band appeared at around 590 nm, as recently reported by Guria et al. [41]. In their approach, a BODIPY probe was designed with a specific architecture bearing a diethyl amino substituted moiety connected to the \textit{meso} position, eventually resulting in a change of the spectral profiles. Few reports following this mechanism have been published for BODIPY probes and amino acid derivatives, and it is possible to find examples with other natural derivative such as G-proteins.

2.2. Molecular Recognition (Non-Covalent)

A different approach for the recognition by BODIPY probes can be used especially for the detection of amino acids and small peptides. For example, precise functional groups can be placed in the BODIPY architecture so that it will specifically interact in a non-covalent fashion with the target molecule, eventually resulting in a change of the spectral profiles. Few reports following this mechanism have been published for BODIPY probes and amino acid derivatives, and it is possible to find examples with other natural derivative such as G-proteins.

One interesting work focused on the rapid detection of aspartic acid and glutamic acid in water was recently reported by Guria et al. [41]. In their approach, a BODIPY probe was designed with a specific architecture bearing a diethyl amino substituted moiety connected to the \textit{meso} position, as shown in Figure 2, 30.

![Scheme 14](image1)

**Scheme 14.** Recognition process for the BODIPY probe 25 toward GSH and proposed molecular displacement for cysteine.

The behaviour of the probe was tested with other bio-thiols such as cysteine and homocysteine. With Cys, after the initial addition of the nucleophile, a new emission band appeared at around 590 nm but rapidly decreased, favouring another band at 564 nm according to the proposed intermolecular displacement shown in Scheme 12 and resulting in the formation of 29. In case of Hcy, similar to GSH, an emission band at 587 nm appeared but it was followed by a strong decrease in the intensity, following the same reactivity as proposed for cysteine.

![Figure 2](image2)

**Figure 2.** The BODIPY sensor that presents a photo-induced electron transfer (PET) process in water.
Because of the specific construction of the probe, a photo-induced electron transfer (PET) is observed with only a very weak emission detectable at 527 nm in water. Upon addition of aspartic acid (Asp) or glutamic acid (Glu) (0–600 μM amino acid, 3:97 DMSO/H2O, pH 6.5), the emission maximum was recorded to rapidly increase as a consequence of the interaction between the probe and the amino acid, which is effectively blocking the PET process. In addition, the absorbance was strongly influenced upon addition of the target molecule; the main absorption band at 514 nm was hypsochromically shifted to 503 nm. The type of interaction between the probe and the target molecule was ascribed to be of hydrogen bonding with theoretical (DFT) and NMR evidence; indeed, significant changes in the chemical shifts of the protons in the pyridine moiety were detected.

The highly selectivity of the system for Asp and Glu was demonstrated by comparing the behaviour with all the other amino acids that, when added to the probe, did not significantly alter the fluorescent profile of the BODIPY. Because of those specific behaviours, the probe was used for intracellular imaging and showed good response and high stability in the cellular systems.

Recently, Guler and co-workers presented an interesting work about a BODIPY probe based on a pillar[5]arene structure with sensing ability for asparagine. The probe (31 in Figure 3) was obtained by click chemistry between an azido BODIPY precursor and a pillar[5]arene having terminal alkynes [42].

![Figure 3. The pillar [5]arene BODIPY sensor presented in Guler’s work.](image)

The cyclic structure presents absorption and emission maxima, respectively, at 325, 505 and 520 nm. Upon addition of thirty different amino acids and derivatives in a DMF/H2O 1:1 mixture, the only appreciable changes were recorded in the emission spectrum when L-asparagine was used. Specifically, the emission intensity was enhanced and a slight blue shift of 6 nm was recorded. The type of interaction between the probe and the sensed specie was described from the authors to be non-covalent with the formation of an inclusion type of complex. The host-guest nature of interaction is supported by NMR experiments, which show how all the protons of the amino acid are reasonably upfield shifted because of the rich electron environment of the cavity.

Kennedy et al. designed and synthetized a GDP analogue integrating a BODIPY moiety (32) (Figure 4) which presents a fluorescence emission with maximum at 510 nm [19].
Interestingly, the incubation of the probe with G-protein’s α subunits in a Tris buffer (192 mM glycine, 1 mM EDTA, 10 mM CaCl₂, pH = 8.5) resulted in a fluorescence anisotropy, indicating the formation of a complex. As control, the probe was tested with trypsin inhibitor and no change in anisotropy was noticeable.

Another example reporting a non-covalent interaction between the probe and the target was documented in 2009 by I. Hamachi and co-workers [20]. In their work, they developed a BODIPY-based probe able to interact and identify the Neurofibrillary Tangles presents in patients affected by the Alzheimer’s disease. They designed a probe with bipyridyl-type Zn(II) complex moieties, i.e., 33, which can easily interact with hyperphosphorylated tau proteins and that presents absorption and emission bands at 520 and 547 nm (Φ = 0.31), respectively (see Figure 5).

Upon addition of the probe in HBS buffer (100 nM) to a suspension of p-Tau (8 μM, overexpressed in Alzheimer’s disease) in HBS buffer containing 10% DMSO, an increase of intensity in the emission was recorded and a dissociation constant of 7.5 μM was calculated. When the sensor was mixed with mono- or non-phosphorylated tau, no changes in the emission could be appreciated, showing a specificity of the probe for the hyperphosphorylated tau proteins.

3. Porphyrin Probes

The structure of porphyrin, namely [18] tetraphyrin(1.1.1.1), provides three types of modification sites: eight β positions, four meso positions and the coordination cavity (Figure 6). Porphyrin hosts for amino acid derivatives typically make use of both the peripheral alterations and metal coordination, the latter being of key importance for such applications. The meso-substituents typically include hydrogen bond donors (e.g., hydroxyl groups) and acceptors (e.g., methoxyl groups) and bulky aromatic rings (e.g., naphthyl). Formation of stable supramolecular complexes is achieved via three-point fixation, consisting of coordination of amino acid N-terminus to the metal centre, as well as hydrogen bonding and electrostatic, hydrophobic or steric interactions.
Porphyrin’s N₄ coordination cavity allows binding of numerous metal ions. A careful choice of metal centre allows coordination of axial ligands, for instance amines [43] or alcohols [44]. Such coordination causes slight perturbation to the optical properties, easily detected with UV/Vis spectroscopy. Examples of sensors described in this section involve zinc(II) and ruthenium(II) porphyrins.

Binding of a chiral molecule to an achiral porphyrin results in chirogenesis-process of transfer of chiral information from the chiral partner (guest) to achiral host [45]. This leads to induction of supramolecular chirality with characteristic strong Cotton effects, carrying information about the guest absolute configuration (Scheme 15) [46]. Formation of supramolecular host-guest complexes described in this paper is therefore easily detected with circular dichroism spectroscopy.

![Figure 6. Porphyrin modification sites.](image)

**Scheme 15.** Schematic presentation of supramolecular chirogenesis in porphyrin-amino acid host-guest systems.

### 3.1. Monomeric Hosts

#### 3.1.1. Early Reports

The use of porphyrins as probes for chiral recognition of amino acids dates back to the early 1990s, when Ogoshi and co-workers published a series of papers describing the design and application of several hosts employing porphyrin zinc complexes with peripheral substituents tailored to bind α-aminoester guests.

Three hosts with various binding capability were examined (Figure 7) as zinc(II) complexes open for coordination of a guest and an auxiliary binding site: a hydrogen bond donor (34) and a hydrogen bond acceptor (35) [47]. Thus, the construction of macrocyclic host was predefined for formation of a doubly bound system with nitrogen coordinated to zinc(II) centre and a hydrogen bond interaction of a C-terminus of an amino acid. The UV/Vis titration with LeuOMe revealed formation of a 1:1 complex 34-LeuOMe upon addition of large excess of the guest (Figure 7) manifested by the Soret band bathochromic shift treated as diagnostic of formation of host-guest system. ¹H NMR experiments
consistently support the participation of hydrogen bonding in host-guest formation. Titration of 34 with a solution of L-LeuOMe results in downfield shift of the naphthyl hydroxy proton signal (marked red in Figure 7) from \( \delta \sim 5.2 \) ppm to \( \delta \sim 6.4 \) ppm, indicating the presence of hydrogen bond incorporating the hydroxyl group. Further experiments performed for methyl esters of valine, phenylalanine, tryptophan, asparagine or glutamic acid showed that bulkier side chains of amino acids positively influence stability of final complexes. In addition, the presence of aromatic substituents increases the stability of final host–guest systems obtained for 35 where hydrogen bond character is inverted while carboxylic groups has no influence. The association constants for 34 (1.7 \( \times \) \( 10^3 \)–1.1 \( \times \) \( 10^4 \) M\(^{-1} \)) are 3.8–40.0 times larger than those obtained for 35. These observations led to the conclusion that hydrogen bonds are formed between naphthyl hydroxyl and aminoester carbonyl groups (Figure 7).

![Figure 7. Ogoshi’s meso-naphthyl sensors for amino acid methyl esters.](image)

Ogoshi and co-workers designed also a condensed \( \alpha,\alpha,\beta,\beta \)-atropoisomer of \( \text{meso-tetra(\text{o-amino})phenyl} \) porphyrin with un- and substituted isophthaloyl chlorides to yield doubly bridged chirality probes 36–39 (Figure 8). Both optically pure enantiomers were examined, and they exhibited significant enantioselectivity, from 4.1 for alanine to 7.5 for valine methyl ester. Upon \(^1\text{H}\) NMR monitored coordination experiment, all guest \(^1\text{H}\) signals shift upfield to negative \( \delta \) values, due to the guest molecule being placed in porphyrin’s magnetic anisotropy cone [48]. In the case of L-ValOMe, the most dramatic shift difference is observed for \( \text{NH}_2 \) and \( \text{C}^\alpha \text{H} \) protons (\( \Delta \delta = -6.04 \) ppm and \( \Delta \delta = -6.21 \) ppm, respectively) as they appear at \( \delta = -4.57 \) ppm and \( \delta = -2.90 \) ppm in the complex, indicating that both groups of protons come in close contact with the porphyrin plane [49].

![Figure 8. Doubly bridged chirality probes.](image)
The O-nitro substituents play an important role in the recognition process. It was found that (-)-35 binds the same guests with association constants 12 and 21 times larger than 37 and 38, indicating that the hydrogen bond donating ability of the amide group remaining in the ortho position to the nitro substituent is increased by the electronic effect of NO₂ group.

3.1.2. Achiral Hosts

Commercially available ruthenium(II) porphyrin complex 40, with a strong Lewis acid character, was employed as a chirality probe for amino acid esters and hydrazides [49]. The strong binding of amines to the Ru centre is reflected in the high association constant determined for coordination of 1-ValOMe as \( K_{\text{ass}} = 1.7 \times 10^7 \) M\(^{-1}\). The difference in coordination abilities to Ru and Zn centres is reflected by the binding constant for 40, which is three-fold higher than those of Zn-based probes, such as 33. Amino acid hydrazides were obtained upon hydrazinolysis of corresponding amino acid methyl esters, and their coordination to 40 was monitored with \(^1\)H NMR. Exposing such hydrazide to 1 eq. of 40 yields two 1:1 complexes, 41 and 42, differing by the donor for coordination to ruthenium. Further addition to the macrocyclic host leads to formation of sandwich complex 43, in which the two terminal amino groups coordinate to ruthenium centres (Figure 9). In such magnetic environment, all \(^1\)H signals of the guest move to negative values of chemical shifts (from \( \delta = -0.5 \) ppm to \( \delta = -6.4 \) ppm). Protons attached to the two terminal amino groups show substantial upfield shifting to \( \delta = -2.8 \) ppm and \( \delta = -3.8 \) ppm (hydrazide NH₂) and \( \delta = -5.4 \) ppm and \( \delta = -6.4 \) ppm (C²-NH₂).

![Figure 9. Stoichiometry-dependent host-guest complex formation.](image)

The presence of chiral guest results in formation of a sandwich complex that exhibits a distinct CD effect in the Soret region (405 and 412 nm). When l isomers were used (valine and phenylalanine), the shorter wavelength band was negative and the other positive, while with d-phenylalanine derivative the CD image was reversed. Some other commercially available ruthenium(II) porphyrins ([RuII(TPP)CO], [RuII(TMP)CO] and [RuII(F20-TPP)CO]) were tested as potential chirality probes and gave corresponding results [49].

3.1.3. Chiral Hosts

Continuing their research on thermodynamic aspects of host-guest interactions between porphyrins and amino acid esters, Ogoshi and co-workers designed a set of chiral hosts based on a zinc(II) coordination centre and auxiliary hydroxyl groups used as hydrogen bond donors (Figure 10). Methyl acetate groups were introduced for a steric repulsion and hydrogen bond
acceptance [50]. While the combination of coordination and hydrogen bonding in the positive isomer gave results corresponding to those reported previously for 35 [47], the presence of ester groups in 44 resulted in slightly repulsive behaviour towards the side chain of L-aminoesters. The effect was even more pronounced in a corresponding series of D-aminoesters. A significant exception was observed for D-SerOBz where the carbonyl group of methyl acetate acted as a hydrogen bond acceptor for serine side chain hydroxyl. Comparing association constants for binding of L- and D-aminoesters, it was concluded that (++)-44 shows significant enantioselectivity for L-aminoesters, except for D-SerOBz, where the enantioselectivity is reversed, probably due to the noteworthy contribution from a second hydrogen bond. The extent of steric repulsion towards L-aminoesters was independent of hydrogen bonding, but in the case of D-aminoesters the repulsion was larger in complexes of 44 (complexes with two-point fixation, i.e., coordination and hydrogen bonding) than 45, where no hydrogen bonding occurs (one-point fixation by coordination). When D isomers were used, the interaction energies were sensitive to the character (size and functional groups) of the guest.

![Figure 10. Three-point fixation of aminoesters in the cavity of porphyrin host.](image1)

2-Hydroxynaphthyl substituent was also employed in the design of a chiral ABCD-type porphyrin reported by Yang et al. Steric hindrance of 2-hydroxyl group and 9-hydrogen atom prevents rotation along the 5–5' bond (Figure 11). Enantioselectivity of (R)-48 was 1.6, favouring L-PheOMe over its chiral counterpart.

![Figure 11. Supramolecular complexes of isomers of ABCD-type porphyrin sensor and L-phenylalanine methyl ester.](image2)

The phenomenon of chiral discrimination of porphyrin-aminoester pairs was employed for NMR-assisted configuration assignment, where phenylalanine methyl esters were used as chiral solvating agents. Formation of host–guest complexes with two-point fixation via NH2-Zn coordination and C=O···HO hydrogen bonding led to decrease in freedom of motion of the host molecules, resulting in resolution of NMR signals in thus obtained diastereoisomeric complexes [51].
3.2. Dimeric Hosts

Due to their 18\(\pi\) electron circuit and ability to accommodate various metal ions inside their cavity, porphyrins have been used as an attractive building block for molecular tweezers [52]. A molecular tweezer is a host molecule consisting of two, usually flat and aromatic subunits, designed to “recognise” specific guest molecules by means of metal coordination, \(\pi-\pi\) interactions, hydrogen bonding, hydrophobic interactions or van der Waals forces. Such subunits are connected via a linker, which can be a spacer of greater or smaller flexibility. Since the first synthesis of a caffeine dimer with diyne linker was reported in 1978 by Chen and Whitlock [53], many molecular tweezers were designed to serve various purposes. Zinc complexes of porphyrins tend to bind neutral ligands through nitrogen, oxygen, phosphorus and sulphur, forming five-coordinated 1:1 adducts [54]. This property has been widely exploited in porphyrin tweezer applications, including those designed for chiral recognition.

In the early 2000s Borovkov et al. reported a chirality induction in achiral ethane-bridged \textit{syn-bis}-porphyrins 49–50 via an interaction with chiral amines and amino acids. Based on the reported dependence of CD signal sign and intensity on the steric bulk of aminoester side chain, they suggested that coordination of the enantiopure ligand induced such a transformation, where the achiral \textit{syn} conformer 49 (Figure 12) effectively transforms into a chiral variant 49\((l\text{-AlaOMe})_2\). In a subsequent study, the mechanism of chirality induction was investigated and tweezers with one or two coordinating centres were employed (Figure 12) [55].

![Figure 12. Achiral ethane-bridged \textit{syn-bis}-porphyrins and their interaction with aminoester guests.](image)

In nonpolar solvents, 49 adapts a cofacial, \textit{syn} conformation, due to strong \(\pi-\pi\) interactions between the two porphyrin rings. Depending on the number of coordination sites, two different \textit{anti}-conformers were observed: 1:2 for two zinc(II) centres and 1:1 for single coordination of zinc(II) as documented spectroscopically.

In UV/Vis, host–guest complexes of 49 and amino acid esters show bathochromic shift of Soret band, as well as its splitting (at 435–437 and 424–426 nm), and a set of two Q bands, among which the higher energy one is significantly more intense than the other. In CD, these systems show strong bisignate couplets in the Soret region—with maxima corresponding to those observed in UV/Vis—and an additional blue-shifted lower intensity Cotton effect.

The factors influencing the handedness of the screw are as follows: right-handed screw is formed when the C-terminus substituent is bulkier than amino acid side chain; in the opposite relation, such arrangement is unfavourable and left-handed screw is induced. Therefore, it was found that the CD response depends not only on the absolute configuration of amino acid guest, but also on the mutual steric relation between the \(C^\alpha\) substituents. Thus, the Cotton effects in the complex only correspond to the guest absolute configuration when the side chain is less bulky than the COOR group.
Among the isomers, positive chirality corresponds to AlaOMe, IleOMe, ValOMe, ProOMe and ProOBn and negative sign was observed in the case of LeuOMe, LeuOBn, PheOMe, PheOBn, PhgOMe, TyrOBn, Tyr(Bn)OMe, Gln(OMe)OMe, GlnOBu, CysOMe, Lys(Z)OMe and TrpOMe.

Investigation of a 1:1 host-guest complex 50-L-AlaOMe (Figure 12) provided insight into the mechanism of chirality induction in such supramolecular complexes. It was demonstrated that steric interaction of 3- and 7'-ethyl groups in 49 and 50 with aminoester guests is the driving force behind chirogenesis (Figure 12), as it locks guest molecule in position. It was also found that the intensity of induced Cotton effects depends on the number of possible chirality-generating conformations, which increases with the number of accommodated aminoester molecules. Moreover, 50 is a suitable host for chiral recognition of bidentate guests, i.e., chiral diamines or amino alcohols, which form 1:1 CD-silent complexes with 49. For example, a 1:1 complex of 49 and 1,2-DACH (1,2-diaminocyclohexane) is so strong, that 1:2 cannot be formed even upon addition of large excess of ligand ($K_{assoc} > 10^7$ M$^{-1}$).

Jiang et al. designed three achiral Zn(II) trip henylporphyrin dimers with amide linkers of various rigidity: 51–53 for the purpose of elucidating the mechanism of chiral recognition in systems with two point fixation. First, oxalic amide-linked tweezer 51 was investigated as a host for nonpolar amino acid ethyl esters. Crystal structure of a 1:2 complex of 51 and rac-PheOEt was solved and the following structural features were brought to light: With a linker of this flexibility, the dimer adapts an anti-configuration in the complex, and both ligand molecules coordinate to zinc from inside the tweezer cavity. Contrary to prior reports, it was found that in 51-(L-PheOEt)(R-PheOEt) hydrogen bonds are formed between aminoester NH$_2$ and linker carbonyl group (Figure 13). For comparison, in their account on meso-naphthyl probe 34, Ogoshi and co-workers suggested that the naphthyl OH group acts as a hydrogen bond donor, and amino acid carbonyl group oxygen is the acceptor (Figure 7) [48].

Figure 13. Porphyrin tweezer with oxalic amide linker and its 1:2 complex with an aminoester guest.

Formation of 1:2 complexes with 51 was documented by UV/Vis titration with solutions of amino acid ethyl esters. Two sets of isosbestic points were detected at 426 and 424 nm. $K_2$ was reported as significantly lower than $K_1$ for all guests employed ($K_1: 2.0–7.2 \times 10^4$ M$^{-1}; K_2: 1.0–1.2 \times 10^4$ M$^{-1}$), consistently with more difficult binding of the second guest due to the different coordination geometry [56]. The intense CD signals of 51-(L-LeuOEt)$_2$ complexes were diagnostic of guest absolute configuration as the accommodation of bulky guests (e.g., LeuOEt) results in induction of strong positive bisignate Cotton effect in the Soret region. The positive Cotton effects induced in host-guest systems indicate that the tweezer adapts a clockwise twist. In such conformation, the aminoester side chain faces outside of the intraporphyrin cavity—the opposite conformer would not be favoured for steric reasons (Figure 13).

In the efforts to provide a more sensitive chirality probe, which would accommodate chiral ligands with higher CD amplitude values, the group designed a bis-porphyrin tweezer with slightly less rigid linker, that is m-phthalic diamide (Figure 14, 52).
Jiang et al. designed three achiral Zn(II) triphenylporphyrin dimers with amide linkers of various rigidity: with two point fixation. First, oxalic amide-linked tweezer 51·(L-PheOEt)(D-PheOEt) hydrogen complexes were diagnostic of guest absolute configuration. Similar results were obtained for an even more flexible malonamide-linked bis-porphyrin tweezer 53 (Figure 14) [58]. Host-guest complexes of this sensor and nonpolar amino acid ethyl esters exhibit CD features closely related to those obtained with a \( m \)-phthalic diamide linked tweezer, namely the sign of induced Cotton effects and their amplitude. Positive sign of induced Cotton effects indicates that tweezers 52 and 53 adapt into a right-handed screw. Hence, it was concluded that chiral recognition is governed by the same principles in those two types of chirality sensors [49,58].

A set of significantly more rigid bis-porphyrin tweezers was reported by Ogoshi and co-workers (Figures 15 and 16) [59]. Porphyrin subunits were connected via chiral 2,2′-dihydroxy- (54) and 2,2′-dimethoxy-1,1′-binaphthy1 (55) as well as achiral 3,3′-biphenyl (56) and 5,5′-dimethoxycarbonyl-3,3′-biphenyl (57) bridges. Such sensors were applied in chiral recognition of lysine derivatives (Figure 15) and cystine diesters (Figure 16), manifested by induction of strong CD response in the Soret region of the host. The bisignate Cotton effects were diagnostic of guest absolute configuration. Association constants for formation of 1:1 complexes were determined by means of UV/Vis spectroscopic titration and remained in the ranges of \( 1.1 \times 10^4 - 1.6 \times 10^5 \) and \( 9.0 \times 10^4 - 2.4 \times 10^6 \) M\(^{-1} \), for binaphthyl- and biphenyl-bridged hosts, respectively.

**Figure 14.** Bis-porphyrin tweezers with \( m \)-phthalic diamide and malonamide linkers.
Flexible tartaric acid-based linker in the design of their sensor for nonpolar amino acid ethyl esters. Molecules 2020, 25, x FOR PEER REVIEW 20 of 28

Figure 15. Binaphthyl-bridged porphyrin tweezers as sensors for lysine derivatives.

Binaphthyl-bridged tweezers 54 and 55 show remarkable enantioselectivity for lysine derivatives, reaching 12 for n/l-Lys-NMe₂, (S)-54 favoring its l isomer (11.5) and (R)-54 its n isomer. It was found that the methoxy groups in the cavity of 54 play a crucial role in chiral differentiation, as corresponding sensor 55 exhibited negligible enantioselectivity towards the same set of guests.

Another example of chiral hosts was recently reported by Wang et al. [60], who used a fairly flexible tartaric acid-based linker in the design of their sensor for nonpolar amino acid ethyl esters.
with a right-handed screw. This chiral sensor shows a negative CD couplet at 416 and 425 nm; coordination of $\text{l}$-amino acid ethyl esters gives rise to a new negative Cotton effect at 426–428, 434 and 436 nm, while their chiral counterparts induce a positive Cotton effect at 424–426 and 430–433 nm. Compound 58 was reported to show enantioselectivity towards $\text{d}$-aminoesters, reaching 8.4, which, according to the authors, is the highest value obtained to date for this class of sensors. Optimised structure of 58 revealed two types of hydrogen bonds—both between linker amino and guest carbonyl groups and the other way around. Peptoids constitute a class of peptidomimetics, with the amino acid side chain attached to the nitrogen instead of the $\alpha$-carbon. In other words, a peptoid is a synthetic $N$-substituted glycine oligomer [61]. Lately, several porphyrin–peptoid conjugates have been designed as energy transfer systems [62]. These hybrids have been of particular interest owing to the simplicity of backbone modification, which is a powerful tool to control the change in inter-chromophore interactions [63,64].

Recently, Seo et al. [65] reported bis-porphyrin tweezers with chiral and achiral peptoids as flexible linkers (Figure 18). Those tweezers were shown to accommodate achiral diamines of various sizes and basicities, but also served the purpose of chiral recognition of lysine stereoisomers, which was monitored with UV/Vis and CD spectroscopies. This bidentate ligand was found to bind to achiral host 59, giving rise to a bisignate Cotton effect in the Soret region (at 425 and 432 nm) reflecting guest absolute configuration. Coordination with $\text{l}$-LysOMe induced a negative Cotton effect, indicating that the complex formed a left-handed screw, while—as expected—$\text{d}$-LysOMe gave a mirror CD image, with a right-handed screw.

When a chiral tweezer 60 was employed, accommodation of chiral guests led to diastereoisomeric supramolecular structures. Upon CD monitored titration of 60 with n-LysOMe, a slight bathochromic shift of the spectrum was observed, from 422 and 430 nm to 425 and 435 nm. A complex with $\text{l}$ isomer gave a diminished “M”-shaped CD response (425, 430 and 435 nm), probably owing to the presence of multiple CD-active conformers in the solution.

(Figure 17). Compound 58 was found to form 1:2 complexes with both $\text{l}$ and $\text{d}$ isomers, with $K_1$ in the range of $4.3 \times 10^3$–$6.8 \times 10^4 \text{ M}^{-1}$ and $K_2$: $6.6 \times 10^2$–$3.0 \times 10^3 \text{ M}^{-1}$. Notably, while initial formation of 1:1 complexes seems to be equally easy as in the case of oxalic amide-linked tweezer reported by Hu et al. [47,56], association constants determined for subsequent coordination of another guest molecule significantly lower in this case. Formation of diastereoisomeric host-guest complexes results in a noticeable bathochromic shift, from 419 nm for 58 to 428 nm for 58·$\text{L}_2$.

![Figure 17. Porphyrin tweezer with tartaric acid-based linker.](image-url)
weakened in water due to hydration [66]. In this paragraph we report some of the recent water-soluble
plane, bound to adjacent meso porphyrin sensors for chiral recognition of amino acids and elucidation of factors influencing such
particular interest, as mimicking physiological conditions provides more precise information useful to
phenomena in biological systems. Therefore, investigation of such interactions in water has been of
61
in the range of 2.4 \times 10^{-10} M^{-1} at pH 7.4 in aqueous solutions are 1–3-fold lower than those reported for organic-phase systems [47] and were
GS, GE, GGG and GGW). Notably, association constants for formation of such host-guest complexes in
amino carboxylates were compared, together with di- and tripeptides (GG, GA, GV , GL, GF, GW, GP ,
60
benzamide moieties were embedded for host-guest hydrophobic attractions. Enantiomers of
improved by Imai et al. [11,67]. Their sensor bears two quaternary ammonium groups above and below porphyrin
by Imai et al. [11,67]. Their sensor bears two quaternary ammonium groups above and below porphyrin
interactions in aqueous media.

Figure 18. Porphyrin tweezers with achiral and chiral peptoid linkers.

3.3. Aqueous Phase

Studies on chiral recognition of biomolecules offer insight into corresponding host-guest phenomena in biological systems. Therefore, investigation of such interactions in water has been of particular interest, as mimicking physiological conditions provides more precise information useful to elucidate specific mechanisms governing chiral recognition in vivo. The vast amount of porphyrin receptors has brought to light some of the most important factors influencing the process of chiral recognition of amino acids in nonpolar solvents, however the environment affects those interactions in a non-negligible way. In organic solutions, hydrogen bonding has been found to play a crucial role in two- (or three-)point fixation of guest molecule onto the porphyrin plane, yet it is considerably weakened in water due to hydration [66]. In this paragraph we report some of the recent water-soluble porphyrin sensors for chiral recognition of amino acids and elucidation of factors influencing such interactions in aqueous media.

Among early reports is a water-soluble porphyrin host for chiral recognition of amino acids reported by Imai et al. [11,67]. Their sensor bears two quaternary ammonium groups above and below porphyrin plane, bound to adjacent meso-phenyl substituents (Figure 19). At the two remaining meso-phenyls, benzamide moieties were embedded for host-guest hydrophobic attractions. Enantiomers of 60 were tested for binding amino acids in water solutions at basic pH values, where their carboxylic group is deprotonated. In this study, charged (Ser, Asp and Glu) and nonpolar (Gly, Ala, Val, Leu, Phe and Trp) amino carboxylates were compared, together with di- and tripeptides (GG, GA, GV, GL, GF, GW, GP, GS, GE, GGG and GGW). Notably, association constants for formation of such host-guest complexes in aqueous solutions are 1–3-fold lower than those reported for organic-phase systems [47] and were in the range of $2.4 \times 10^{-1} – 1.5 \times 10^{3}$ M$^{-1}$. In aqueous solution, guest molecules compete with H$_2$O coordinated to the zinc centre. Compound 61 showed slight enantioselectivity towards enantiopure ligands, ranging from 1.2 to 3.3.
Coordination of guest NH$_2$ to host zinc(II) centre and Coulomb interactions between cationic meso substituents and deprotonated carboxy groups of the amino acids were found to contribute to the binding constant together with hydrophobic attraction between amino acid side chain and the benzamide phenyl ring or steric repulsion between them. The latter depends on the structure of amino carboxylate in the following way. Among the most tightly bound guests are tryptophan and phenylalanine; less hydrophobic residues, such as serine, aspartic acid or glycylerine, do not show an increase in association constant, yet their stereoisomers are bound to the host selectively. Dependence of association constant on ionic strength of the solution was determined, supporting the contribution of electrostatic interactions to formation of host-guest complexes (n-butylamine was used as reference). Recognition ability towards GX dipeptides (where X = G, A, V, L, F, W, P, S or E) was retained, even though the asymmetric centre is separated from the coordination site. Chiral recognition of nonpolar amino acids by ruthenium Halterman porphyrins in chloroform, methanol and water was reported by Nicolas et al. [68]. In this study, a rigid porphyrin sensor 62 with Ru(CO) centre was employed to investigate ligation of the metal centre in trans position to the other axial ligand (Figure 20).

The application of a chiral host for recognition enabled the differentiation and quantification of diastereoisomeric complexes formed upon addition of racemic ligands in $^1$H NMR. Among the chiral
amines investigated were alanine, phenylalanine and valine as acids (as guests for 63 in aqueous phase) and methyl esters (for 62 in organic phase studies). In all environments, \( \text{n-phenylalanine} \) was favoured over its enantiomeric counterpart, while Val and Ala showed the opposite preference. In the case of phenylalanine, a drastic decrease of enantioselectivity was observed while changing the environment from organic to aqueous; it was therefore concluded that hydrophobic interactions play a significant but not crucial role in the process of chiral recognition.

Based on previous studies on the origin of chiral recognition and its dependence on different association and dissociation constants in the formed complexes [69,70], the latter were examined with saturation transfer NMR technique, exploiting the slow axial ligand exchange in ruthenium porphyrins [45]. The data obtained from these experiments reveal that dissociation rate constants were lower for isomers, towards which selectivity was observed (i.e., \( k_{d-PheOMe} = 0.36 \text{ s}^{-1} \) and \( k_{d-PheOMe} = 0.14 \text{ s}^{-1} \)), indicating that the rate of ligand dissociation affects the process of chiral recognition.

4. Conclusions

We highlight the importance of the detection of natural derivatives, specifically focusing on amino acids, by two different molecular probes, BODIPY and porphyrins. The two probes essentially work with two different recognition approaches: BODIPY probes rely on the direct interaction with the target, hence a covalent interaction. Porphyrins probes, on the other hand, mostly interact with the target in the fashion of molecular recognition, which is mediated by non-covalent interactions—a synthesis of coordination, hydrogen bonding, hydrophobic interactions, steric repulsion and/or electrostatic attraction. To this day, sensors for both molecular recognition and chiral discrimination are designed. The development of such systems is of significant importance for the scientific community since there are already many specific applications, especially in the fields of biomedicine and molecular devices.

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References

1. Persch, E.; Dumele, O.; Diederich, F. Molecular Recognition in Chemical and Biological Systems. *Angew. Chem. Int. Ed.* 2015, 54, 3290–3327. [CrossRef] [PubMed]
2. Baron, R.; McCammon, J.A. Molecular Recognition and Ligand Association. *Annu. Rev. Phys. Chem.* 2013, 64, 151–175. [CrossRef] [PubMed]
3. Ariga, K.; Ito, H.; Hillab, J.P.; Tsukube, H. Molecular recognition: From solution science to nano/materials technology. *Chem. Soc. Rev.* 2012, 41, 5800–5835. [CrossRef] [PubMed]
4. Nakamura, T.; Kaneko, Y.; Nishibori, E.; Nabeshima, T. Molecular Recognition by Multiple Metal Coordination inside Wavy-Stacked Macrocycles. *Nat. Commun.* 2017, 8, 1–7. [CrossRef]
5. Fan, E.; Van Arman, S.A.; Kincaid, S.; Hamilton, A.D. Molecular Recognition: Hydrogen-Bonding Receptors That Function in Highly Competitive Solvents. *J. Am. Chem. Soc.* 1993, 115, 369–370. [CrossRef]
6. Yao, H.; Ke, H.; Zhang, X.; Pan, S.-J.; Li, M.-S.; Yang, L.-P.; Schreckenbach, G.; Jiang, W. Molecular Recognition of Hydrophilic Molecules in Water by Combining the Hydrophobic Effect with Hydrogen Bonding. *J. Am. Chem. Soc.* 2018, 140, 13466–13477. [CrossRef]
7. Yang, L.; Adam, C.; Nicol, G.S.; Cockroft, S.L. How Much Do van Der Waals Dispersion Forces Contribute to Molecular Recognition in Solution? *Nat. Chem.* 2013, 5, 1006–1010. [CrossRef]
8. Mueldorph, A.V.; Van Engen, D.; Warner, J.C.; Hamilton, A.D. Aromatic-Aromatic Interactions in Molecular Recognition: A Family of Artificial Receptors for Thymine That Shows Both Face-to-Face and Edge-to-Face Orientations. *J. Am. Chem. Soc.* 1988, 110, 6561–6562. [CrossRef]
9. Crowley, J.D.; Goshe, A.J.; Bosnich, B. Molecular Recognition. Electrostatic Effects in Supramolecular Self-Assembly. *Chem. Commun.* 2003, 9, 392–393. [CrossRef]
10. Hossoowi, L.; Hong, K.I.; Jang, W.D. Design and Applications of Molecular Probes Containing Porphyrin Derivatives. *Coord. Chem. Rev.* 2018, 354, 46–73. [CrossRef]
11. Imai, H.; Munakata, H.; Uemori, Y.; Sakura, N. Chiral Recognition of Amino Acids and Dipeptides by a Water-Soluble Zinc Porphyrin. *Inorg. Chem.* **2004**, *43*, 1211–1215. [CrossRef]

12. Randazzo, R.; Gaeta, M.; Gangemi, C.M.A.; Fragalà, M.E.; Purrello, R.; D’Urso, A. Chiral Recognition of α- and δ-Amino Acid by Porphyrin Supramolecular Aggregates. *Molecules* **2019**, *24*, 84. [CrossRef]

13. Huang, X.; Nakanishi, K.; Berova, N. Porphyrins and Metalloporphyrins: Versatile Circular Dichroic Reporter Groups for Structural Studies. *Chirality* **2000**, *12*, 237–255. [CrossRef]

14. Davis, S.J.; Ikekizu, S.; Evans, E.J.; Fugger, L.; Bakker, T.R.; van der Merwe, P.A. The Nature of Molecular Recognition by T Cells. *Nat. Immunol.* **2003**, *4*, 217–224. [CrossRef] [PubMed]

15. Weerapana, E.; Wang, C.; Simon, G.M.; Richter, F.; Khare, S.; Dillon, M.B.D.; Bachovchin, D.A.; Mowen, K.; Baker, D.; Cravatt, B.F. Quantitative reactivity profiling predicts functional cysteine in proteomes. *Nature* **2010**, *468*, 790–795. [CrossRef] [PubMed]

16. Zhou, Y.; Yoon, J. Recent progress in fluorescent and colorimetric chemosensors for detection of amino acids. *Chem. Soc. Rev.* **2012**, *41*, 52–67. [CrossRef]

17. Cheng, Z.; Kuru, E.; Sachdeva, A.; Vendrell, M. Fluorescent amino acids as versatile building blocks for chemical biology. *Nat. Rev. Chem.* **2020**, *4*, 275–290. [CrossRef]

18. Kaur, P.; Singh, K. Recent Advances in the Application of BODIPY in Bioimaging and Chemosensing. *J. Mater. Chem. C* **2019**, *7*, 11361–11405. [CrossRef]

19. Jameson, E.E.; Cunliffe, J.M.; Neubig, R.R.; Sunahara, R.K.; Kennedy, R.T. Detection of G Proteins by Affinity Probe Capillary Electrophoresis Using a Fluorescently Labeled GTP Analogue. *Anal. Chem.* **2003**, *75*, 4297–4304. [CrossRef]

20. Ojida, A.; Sakamoto, T.; Inoue, M.; Fujishima, S.; Lippens, G.; Hamachi, I. Fluorescent BODIPY-Based Zn(II) Complex as a Molecular Probe for Selective Detection of Neurofibrillar Tangles in the Brains of Alzheimer’s Disease Patients. *J. Am. Chem. Soc.* **2009**, *131*, 6543–6548. [CrossRef]

21. Zhang, J.; Wang, N.; Ji, X.; Tao, Y.; Wang, J.; Zah, W. BODIPY-Based Fluorescent Probes for Biothiols. *Chem. Eur. J.* **2020**, *26*, 4172–4192. [CrossRef] [PubMed]

22. Farinone, M.; Czybińska, J.; Pawlicki, M. A Controlled Blue-Shift in Meso-Nitrogen Aryl Fused DIPY and BODIPY Skeletons. *Org. Chem. Front.* **2019**, *6*, 2852–2852. [CrossRef]

23. Farinone, M.; Czybinska, J.; Pawlicki, M. BODIPY-Amino Acid Conjugates—Tuning the Optical Response with a Meso-Heteroatom. *Org. Chem. Front.* **2020**, *7*, 2391–2398. [CrossRef]

24. Sørensen, M.L.H.; Vosch, T.; Laursen, B.W.; Hansen, T. Spectral Shifts of BODIPY Derivatives: A Simple Continuous Model. *Photochem. Photobiol. Sci.* **2019**, *18*, 1315–1323. [CrossRef] [PubMed]

25. Loundet, A.; Burgess, K. BODIPY Dyes and Their Derivatives: Syntheses and Spectroscopic Properties. *Chem. Rev.* **2007**, *107*, 4891–4932. [CrossRef]

26. Bañuelos, J. BODIPY Dye, the Most Versatile Fluorophore Ever? *Chem. Rec.* **2016**, *16*, 335–348. [CrossRef]

27. Prasannan, D.; Arunkumar, C. A “Turn-on-and-off” PH Sensitive BODIPY Fluorescent Probe for Imaging *E. Coli* Cells. *New J. Chem.* **2018**, *42*, 3473–3482. [CrossRef]

28. Bricks, J.L.; Kovalchuk, A.; Trießinger, C.; Noz, M.; Bueschel, M.; Tolmachev, A.I.; Daub, J.; Rurack, K. On the Development of Sensor Molecules that Display FIII-amplified Fluorescence. *J. Am. Chem. Soc.* **2005**, *127*, 13522–13529. [CrossRef]

29. Zhang, J.; Pana, F.; Jina, Y.; Wang, N.; Hea, J.; Zhanga, W.; Zhao, W. A BODIPY-based dual-responsive turn-on fluorescent probe for NO and nitrite. *Dyes Pigment.* **2018**, *155*, 276–283. [CrossRef]

30. Shao, J.; Guo, H.; Ji, S.; Zhao, J. Styryl-BODIPY Based Red-Emitting Fluorescent OFF–ON Molecular Probe for Specific Detection of Cysteine. *Biosens. Bioelectron.* **2011**, *26*, 3012–3017. [CrossRef]

31. Wang, C.; Xia, X.; Luo, J.; Qian, Y. A Novel Near-Infrared Styryl-BODIPY Fluorescent Probe for Discrimination of GSH and Its Application in Living Cells. *Dyes Pigment.* **2018**, *152*, 85–92. [CrossRef]

32. Wang, Q.; Wei, X.; Li, C.; Xie, Y. A Novel P-Aminophenylthio- and Cyano-Substituted BODIPY as a Fluorescence Turn-on Probe for Distinguishing Cysteine and Homocysteine from Glutathione. *Dyes Pigment.* **2018**, *148*, 212–218. [CrossRef]

33. Leen, V.; Yuan, P.; Wang, L.; Boers, N.; Dehaen, W. Synthesis of Meso-Halogenated BODIPYs and Access to Meso-Substituted Analogues. *Org. Lett.* **2012**, *14*, 6150–6153. [CrossRef]

34. Jeon, S.; Kim, T.; Jin, H.; Lee, U.; Bae, J.; Bouffard, J.; Kim, Y. Amine- Reactive Activated Esters of Meso-CarboxyBODIPY: Fluorogenic Assays and Labeling of Amines, Amino Acids, and Proteins. *J. Am. Chem. Soc.* **2020**, *142*, 9231–9239. [CrossRef] [PubMed]
35. Jiang, X.; Yue, S.; Jia, L.; Li, S.; Li, C.; Li, Q.; Xiao, L. NIR Fluorescent AzaBODIPY-Based Probe for the Specific Detection of L-Lysine. Chem. Sel. 2018, 3, 7581–7585. [CrossRef]

36. Adhikari, S.; Ghosh, A.; Mandal, S.; Guria, S.; Banerjee, P.P.; Chatterjee, A.; Das, D. Colorimetric and fluorescence probe for the detection of nano-molar lysine in aqueous medium. Org. Biomol. Chem. 2016, 14, 10688–10694. [CrossRef]

37. Liu, Y.; Lv, X.; Hou, M.; Shi, Y.; Guo, W. Selective Fluorescence Detection of Cysteine over Homocysteine and Glutathione Based on a Cysteine-Triggered Dual Michael Addition/Retro-Aza-Aldol Cascade Reaction. Anal. Chem. 2015, 87, 11475–11483. [CrossRef]

38. Avellanal-Zaballa, E.; Ramos-Torres, A.; Prieto-Castañeda, A.; García-Garrido, F.; Bañuelos, J.; Agararbeitia, A.R.; Ortiz, M.J. A BODIPY-Based Fluorescent Sensor for Amino Acids Bearing Thiol. Proceedings 2019, 41, 18. [CrossRef]

39. Wang, N.; Wang, Y.; Gao, J.; Ji, X.; He, J.; Zhang, J.; Zhao, W. A Ratiometric Fluorescent BODIPY-Based Probe for Rapid and Highly Sensitive Detection of Cysteine in Human Plasma. Analyst 2018, 143, 5728–5735. [CrossRef]

40. Niu, L.; Guan, Y.; Chen, Y.; Wu, L.; Tung, C.; Yang, Q. BODIPY-Based Ratiometric Fluorescent Sensor for Highly Selective Detection of Glutathione over Cysteine and Homocysteine. J. Am. Chem. Soc. 2012, 134, 18928–18931. [CrossRef]

41. Guria, S.; Ghosh, A.; Manna, K.; Pal, A.; Adhikary, A.; Adhikari, S. Rapid detection of aspartic acid in water by BODIPY-Based fluorescent probe: Live-cell imaging and DFT studies. Dyes Pigment. 2019, 168, 111–122. [CrossRef]

42. Bastug, E.; Kursunlu, A.N.; Guler, E. A fluorescent clever macrocycle: Deca-bodipy bearing a pillar [5]arene and its selective binding of asparagine in half-aqueous medium. J. Lumin 2020, 225, 117343. [CrossRef]

43. Mizutani, T.; Wada, K.; Kitagawa, S. Molecular Recognition of Amines and Amino Esters by Zinc Porphyrin Receptors: Binding Mechanisms and Solvent Effects. J. Org. Chem. 2000, 65, 6097–6106. [CrossRef] [PubMed]

44. Hayashi, S.; Yotsukura, M.; Noji, M.; Takanami, T. Bis(Zinc Porphyrin) as a CD-Sensitive Bidentate Host Molecule: Direct Determination of Absolute Configuration of Mono-Alcohols. Chem. Commun. 2015, 51, 11068–11071. [CrossRef] [PubMed]

45. Borovkov, V.V.; Lintuluoto, J.M.; Inoue, Y. Supramolecular Chirogenesis in Zinc Porphyrins: Mechanism, Role of Guest Structure, and Application for the Absolute Configuration Determination. J. Am. Chem. Soc. 2001, 123, 2979–2989. [CrossRef]

46. Allenmark, S. Induced Circular Dichroism by Chiral Molecular Interaction. Chirality 2003, 15, 409–422. [CrossRef]

47. Mizutani, T.; Ema, T.; Yoshida, T.; Kuroda, Y.; Ogoshi, H. Recognition of Alpha-Amino Acid Esters by Zinc Porphyrin Derivatives via Coordination and Hydrogen Bonding Interactions. Evidence for Two-Point Fixation from Thermodynamic and Induced Circular Dichroism Spectroscopic Studies. Inorg. Chem. 1993, 32, 2072–2077. [CrossRef]

48. Kuroda, Y.; Kato, Y.; Higashioji, T.; Hasegawa, J.; Kawanami, S.; Takahashi, M.; Shiraishi, N.; Tanabe, K.; Ogoshi, H. Chiral Amino Acid Recognition by a Porphyrin-Based Artificial Receptor. J. Am. Chem. Soc. 1995, 117, 10950–10958. [CrossRef]

49. Liang, Q.-F.; Liu, J.-J.; Chen, J. Sandwich Structure of a Ruthenium Porphyrin and an Amino Acid Hydrazide for Probing Molecular Chirality by Circular Dichroism. Tetrahedron Lett. 2011, 52, 3987–3991. [CrossRef]

50. Mizutani, T.; Ema, T.; Tomita, T.; Kuroda, Y.; Ogoshi, H. Design and Synthesis of a Trifunctional Chiral Porphyrin with C2 Symmetry as a Chiral Recognition Host for Amino Acid Esters. J. Am. Chem. Soc. 1994, 116, 4204–4250. [CrossRef]

51. Yang, L.; Zhou, Y.; Zhu, M.; Zhao, L.; Wei, L.; Bian, Y. Stereochemistry and Solid-State Structure of an Intrinsically Chiral Meso-Patterned Porphyrin: Case Study by NMR and Single-Crystal X-Ray Diffraction Analysis. J. Org. Chem. 2013, 78, 9949–9955. [CrossRef]

52. Valderrey, V.; Aragay, G.; Ballester, P. Porphyrin Tweezer Receptors: Binding Studies, Conformational Properties and Applications. Coord. Chem. Rev. 2014; 258, 259, 137–156. [CrossRef]

53. Chen, C.W.; Whitlock, H.W. Molecular Tweezers: A Simple Model of Bifunctional Intercalation. J. Am. Chem. Soc. 1978, 100, 4921–4922. [CrossRef]
54. Sanders, J.K.M.; Bampos, N.; Clyde-Watson, Z.E.; Darling, S.L.; Hawley, J.C.; Kim, H.-J.; Mak, C.C.; Webb, S.J. Axial Coordination Chemistry of Metalloporphyrins. In The Porphyrin Handbook; Kadish, K.M., Smith, K.M., Guilard, R., Eds.; Academic Press: London, UK, 2000; Volume 3, pp. 30–31.

55. Borovkov, V.V.; Hembury, G.A.; Yamamoto, N.; Inoue, Y. Supramolecular Chirogenesis in Zinc Porphyrins: Investigation of Zinc-Freebase Bis-Porphyrin, New Mechanistic Insights, Extension of Sensing Abilities, and Solvent Effect. J. Phys. Chem. A 2003, 107, 8677–8686. [CrossRef]

56. Jiang, J.; Feng, Z.; Liu, B.; Hu, C.; Wang, Y. Chiral Recognition of Amino Acid Esters by a Novel Oxalic Amide-Linked Bisporphyrin. Dalton Trans. 2013, 42, 7651–7659. [CrossRef] [PubMed]

57. Jiang, J.; Fang, X.; Liu, B.; Hu, C. M-Phthalic Diamide-Linked Zinc Bisporphyrinate: Spontaneous Resolution of Its Crystals and Its Application in Chiral Recognition of Amino Acid Esters. Inorg. Chem. 2014, 53, 3298–3306. [CrossRef] [PubMed]

58. Liu, B.; Jiang, J.; Fang, X.; Hu, C. Absolute Configurational Assignments of Amino Acid Esters by a CD-Sensitive Malonamide-Linked Zinc Bisporphyrinate Host. Chin. J. Chem. 2014, 32, 797–802. [CrossRef]

59. Hayashi, T.; Aya, T.; Nonoguchi, M.; Mizutani, T.; Hisaeda, Y.; Kitagawa, S.; Ogoshi, H. Chiral Recognition and Chiral Sensing Using Zinc Porphyrin Dimers. Tetrahedron 2002, 58, 2803–2811. [CrossRef]

60. Wang, J.; Zhang, Z.; Hu, C.; Wang, Y. Enantioselectivity of a Tartaric Acid Amide Linked Zinc Bisporphyrinate towards Amino Acid Esters. Dyes Pigment. 2020, 176, 108223. [CrossRef] [PubMed]

61. Zuckermann, R.N.; Kerr, J.M.; Kent, S.B.H.; Moos, W.H. Efficient Method for the Preparation of Peptoids by Submonomer Solid-Phase Synthesis. J. Am. Chem. Soc. 1992, 114, 10646–10647. [CrossRef]

62. Kang, B.; Yang, W.; Lee, S.; Mukherjee, S.; Forstater, J.; Kim, H.; Goh, B.; Kim, T.-Y.; Voelz, V.A.; Pang, Y.; et al. Precisely Tuneable Energy Transfer System Using Peptoid Helix-Based Molecular Scaffold. Sci. Rep. 2017, 7, 1–10. [CrossRef]

63. Yang, W.; Kang, B.; Voelz, V.A.; Seo, J. Control of Porphyrin Interactions via Structural Changes of a Peptoid Scaffold. Org. Biomol. Chem. 2017, 15, 9670–9679. [CrossRef] [PubMed]

64. Kang, B.; Chung, S.; Ahn, Y.D.; Lee, J.; Seo, J. Porphyrin–Peptoid Conjugates: Face-to-Face Display of Porphyrins on Peptoid Helices. Org. Lett. 2013, 15, 1670–1673. [CrossRef] [PubMed]

65. Lee, Y.J.; Kang, B.; Seo, J. Metalloporphyrin Dimers Bridged by a Peptoid Helix: Host-Guest Interaction and Chiral Recognition. Molecules 2018, 23, 2741. [CrossRef] [PubMed]

66. Mizutani, T.; Wada, K.; Kitagawa, S. Porphyrin Receptors for Amines, Amino Acids, and Oligopeptides in Water. J. Am. Chem. Soc. 1999, 121, 11425–11431. [CrossRef]

67. Imai, H.; Misawa, K.; Munakata, H.; Uemori, Y. Water-Soluble Zinc Porphyrins as Receptors for Amino Carboxylates. Chem. Lett. 2001, 30, 688–689. [CrossRef]

68. Nicolas, I.; Chevance, S.; Maux, P.L.; Simonneaux, G. Chiral Recognition of Amines and Amino Acid Derivatives by Optically Active Ruthenium Halterman Porphyrins in Organic Solvents and Water. Tetrahedron Asymmetry 2010, 21, 1788–1792. [CrossRef]

69. Maux, P.L.; Bahri, H.; Simonneaux, G. Molecular Recognition of Racemic Phosphines by a Chiral Ruthenium Porphyrin. J. Chem. Soc. Chem. Commun. 1991, 1350–1352. [CrossRef]

70. Galardon, E.; Le Maux, P.; Bondon, A.; Simonneaux, G. Chiral Recognition of Amino Esters by a Ruthenium Porphyrin Complex: Kinetics of the Exchange Process Determined by $^1$H NMR. Tetrahedron Asymmetry 1999, 10, 4203–4210. [CrossRef]