

REVIEW

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Illuminating microtubule functions with small molecules: past, present and future of fluorescent tubulin-binding probes

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Developing tools for the real-time visualization of tubulin, microtubules, and their dynamics is essential for studying key cellular processes such as cell division, migration, and differentiation. In this context, fluorescent small-molecule probes that bind to distinct and specific sites on microtubules have significantly advanced our understanding of tubulin dynamics and functions. In addition, fluorescent probes targeting tubulin have enabled the development of high-throughput screening assays to identify novel anti-tubulin compounds. In this review, we present a comprehensive overview of currently available fluorescent molecules that bind to tubulin, detailing their chemical structures, binding sites, development strategies and relevance to the study of microtubule functions. We also explore future directions in this field and highlight key fluorescent probes that are still lacking. Developing these tools would enable the investigation of tubulin's diverse cellular roles in even greater depth and support the advancement of tubulin-targeted therapeutics.

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Introduction

Microtubules are cytoskeletal polymers formed by the polarized assembly of α/β -tubulin heterodimers, and their behavior is tightly regulated at multiple levels. Through their dynamic properties, microtubules perform essential cellular functions including ensuring proper chromosome segregation during mitosis, facilitating intracellular transport *via* motor proteins, forming specialized motile structures such as cilia and flagella, and enabling cells to acquire and maintain their specific shapes. Given their central role in cell division, shaping and angiogenesis, microtubules are crucial for cell proliferation and migration (extensively reviewed in a previous study¹), one of the

hallmarks of cancer, along with the need for vascular support. Consequently, microtubule-targeting agents (MTAs) have been among the most successful classes of drugs used in cancer therapy (reviewed in previous works^{2–4}). These compounds are broadly categorized as microtubule stabilizers (MSAs) or destabilizers (MDAs) (including tubulin degradation inducers) based on their overall effect on the equilibrium between soluble tubulin and microtubule polymers. To date, eight distinct ligand-binding sites on tubulin have been described, allowing MTAs to be classified by their specific binding domain.^{5–7} Named after the first compound discovered to bind to each specific region, these druggable sites in tubulin are the colchicine site (1), the vinca domain (2), the taxane binding site (3), the maytansine binding site (4), the laulimalide/peloruside binding site (5), the pironetin binding site (6), the gatorbulin binding site (7), and the todalam binding site (8) (Fig. 1). Despite their efficacy, MTAs often have limited clinical use due to their adverse effects, particularly neurotoxicity and myelosuppression, which hinder their broader applications in other tubulin-related pathologies (reviewed in previous studies^{8–10}). Hence, a better understanding of the action of MTAs at the molecular, cellular, and organismal levels is essential for developing safer and more effective compounds.

Historically, studying microtubules and the effects of small molecules on their dynamics has required a multidisciplinary approach. The discovery of tubulin itself was enabled by its interactions with colchicine, which led to tubulin purification and allowed biochemical characterization of microtubules.^{11,12} The development of tubulin-specific antibodies allowed the visualization of the cytoskeleton and the evaluation of drug effects on microtubule's functions.^{13–16} However, this method requires cell fixation and permeabilization, thus limiting dynamic studies. Early live-cell imaging of microtubule behavior relied on microinjecting a rhodamine-labeled tubulin,



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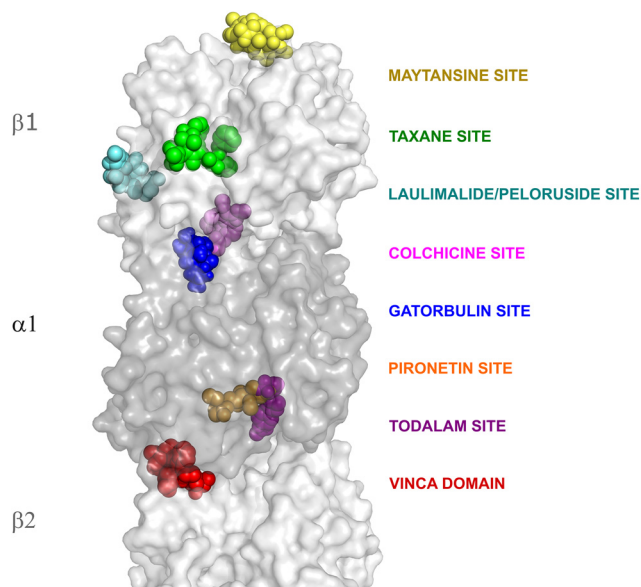


Fig. 1 Binding sites described for small-molecule ligands that can interfere with microtubule functions. Space-filling model of a representative ligand for each binding site is shown in different colors, with the site name in the same color. α -tubulin is shown in dark grey and β -tubulin in light grey.

a technically challenging method requiring tubulin isolation, labeling and reintroduction into cells.^{17–19} A major breakthrough came in the late 1990s with the introduction of plasmids encoding green fluorescent protein (GFP)-tagged cytoskeletal proteins into eukaryotic cells (reviewed in a previous study²⁰), leading to the first cell lines stably expressing GFP- α -tubulin.²¹ However, while the impact of this technique on the field has been significant, overexpression and tag placement can alter the tubulin isotype composition and microtubule dynamics, which limits the interpretability of results obtained with these systems.^{21–23} To overcome these shortcomings, more recent studies have used fluorescent tags embedded within the tubulin sequence to preserve the native isotype expression.²⁴ Moreover, gene-editing technologies now enable endogenous tagging of tubulin, avoiding disturbances in tubulin homeostasis and allowing the implementation of single-molecule imaging.^{25–27}

An alternative approach has been fusing fluorescent proteins to microtubule-associated proteins (MAPs) for indirect microtubule visualization and simultaneous study of MAP-microtubule interactions both biochemically and in cells.^{28,29} Recently, fluorescent peptides derived from MAPs have been developed to probe microtubule functions with minimal perturbation.³⁰

Furthermore, alternative peptide-based approaches have encompassed the identification of short peptide sequences *via* phage display, in which the authors conjugated them to different fluorophores to suit specific experimental requirements.³¹

In addition to protein-based approaches, biologically active fluorescent probes derived from tubulin-binding drugs or other ligands have proven to be effective for visualizing and studying microtubules. Although these compounds must be carefully

selected and dosed to avoid interference with microtubule dynamics, they offer significant advantages. Their versatile synthetic accessibility and increasingly improving photophysical properties provide a high signal-to-noise ratio that enables single-molecule sensing. The conjugation of low molecular weight fluorophores to target-specific recognition elements enables the labeling of molecular structures with extraordinary levels of precision. Furthermore, low molecular weight probes provide access to the nanometer scale, thus allowing to reach confined environments that could remain inaccessible to macromolecular options. Finally, fluorescent tubulin probes can be used for the *in vitro* screening and characterization of novel compounds, particularly to determine drug-binding sites *via* competition assays based on the changes in fluorescence signal. It is also worth pointing out that some compounds exhibit intrinsic fluorescence upon binding to tubulin, enabling straightforward and precise affinity measurements.³²

Although ligand-derived molecular scaffolds provide excellent templates for fluorescent tubulin probes, their development remains a challenge due to synthetic and biochemical hurdles. However, major methodological advances have been reported in the field in recent years, which are summarized in this review together with their impact on cytoskeletal research. Fluorescent probes will be grouped together according to their binding site on tubulin, which will be discussed following the chronological order of their discovery. In addition to reviewing the data on tubulin-directed fluorescent probes reported in the relevant literature, we report previously unpublished data from our own laboratory, including hitherto unavailable equilibrium-binding constants (K_b) for several commercially available fluorescent tubulin probes. We believe that binding constants are essential parameters to be considered when choosing an appropriate probe. Finally, our account includes a tabulated summary of the most relevant probes, detailing their chemical structures, binding affinities (where available), and reported applications. This data collection is intended to support the informed selection of molecular tools for experimental design in tubulin/microtubule research.

It should be noted here that, in addition to fluorescent probes based on small molecules that target the well-characterized binding sites in tubulin, our analysis of the literature has identified other compounds reported to associate with tubulin. For example, DAPI, a DNA intercalator commonly used for nuclear staining (reviewed in a previous study³³), has also been shown to bind to tubulin,³⁴ and it has been employed in *in vitro* studies of polymerization dynamics.³⁵ Similarly, other fluorescent compounds such as bis-ANS,³⁶ DCVJ,³⁷ Nile red,³⁸ and prodan³⁹ have been shown to interact with tubulin and have been used in various *in vitro* experimental settings. However, these compounds are neither MSAs nor MDAs, and their binding to tubulin is not specific. For these reasons, they will not be further discussed in this review.

Colchicine site

Colchicine is a microtubule destabilizer and the first microtubule-targeting agent ever identified. In fact, colchicine



enabled the discovery of tubulin as the main structural component of microtubules by Borisy and Taylor in 1967.^{40,41} They characterized tubulin as a protein capable of binding to colchicine and concluded that this protein served as the fundamental building block of microtubules. Although numerous compounds have since been identified to bind to the colchicine site, none of these have proven suitable for cancer clinical applications, due to their high toxicity and narrow therapeutic window.⁴² However, colchicine itself is currently used for the treatment of gout and various autoimmune diseases. Additionally, some azoles such as mebendazole and albendazole are employed as anthelmintics (reviewed in previous studies^{43,44}). Finally, podophyllotoxin has been widely used in the topical treatment of genital warts^{45,46} and infections caused by *Molluscum contagiosum* virus.^{47,48}

Structurally, the colchicine site is located within a sizable hydrophobic pocket at the interface of the α - and β -tubulin subunits within the same heterodimer. This pocket is defined by helices H7 and H8, the T7 loop, and strands S8, S9 and S10 of the two tubulin β -sheets, and it is completed by the T5 loop of α -tubulin⁴⁹ (Fig. 2). The compound's accommodation in this site generates several conformational adjustments that explain its inhibitory activity. The most significant of these is a switch in the T7 loop that flips to accommodate the ligand, thus blocking the curved-to-straight transition necessary for microtubule assembly, thereby inhibiting polymerization.^{50,51} The colchicine binding site can be subdivided into three zones.⁵² Zone 2 serves as the primary binding pocket and accommodates most ligand structures. Zones 1 and 3 are auxiliary, with very few ligands occupying both simultaneously. Zone 1 lies at the interface with the α -subunit, while zone 3 extends deeper into the β -subunit. Ligands with a globular or butterfly-like conformation, referred to as classical colchicine site inhibitors (CBSIs), typically interact with zones 1 and 2, mimicking colchicine's binding mode. In contrast, non-classical CBSIs,

which exhibit more planar structures, tend to bind deeper within the β -subunit, covering zones 2 and 3.^{53–57}

Colchicine (**1a**) itself fluoresces upon binding to tubulin, and therefore, it is described as the first fluorogenic probe to target this protein. This property was first reported by Arai and Okuyama in 1973^{58,59} and independently confirmed by Bhattacharyya and Wolff in 1974.⁶⁰ This feature allowed the biochemical determination of the behavior of this ligand when interacting with tubulin. However, colchicine presents a low kinetic dissociation constant and almost irreversibly binds to tubulin, which limits its applicability for the characterization of novel compounds in competitive experiments. To address this shortcoming, a new fluorescent compound, 2-methoxy-5-(2',3',4'-trimethoxyphenyl) tropone (also known as MTC, **1b**) was developed by Fitzgerald.⁶¹ This compound, lacking the ring B of colchicine, retains the fluorescence properties of the parent compound but exhibits faster reversible binding⁶² (K_b of $4.9 \times 10^5 \text{ M}^{-1}$).⁶³ This probe has been widely employed to characterize both the colchicine site and its binders, allowing the identification and characterization of many novel ligands of this domain.^{64–69}

As an alternative to exploiting the intrinsic fluorescence of colchicine and derivatives, different fluorophores have been attached to the molecule to allow a broader use *in cells* and *ex vivo* scenarios (Fig. 3). In 1978, Clark and Garland attached a fluorescein molecule at C7,⁷⁰ resulting in a probe (**1c**) that showed similar biological behavior as the parent compound. However, in 1999, Craenenbroeck and Engelborghs demonstrated that the fluorescein group altered the binding behavior, thereby identifying a secondary colchicine-binding site and classifying the probe as a conformation-dependent binder.⁷¹ Despite this limitation, the fluorescein-colchicine probe has been extensively used to characterize colchicine analogues.^{72–74} In 2018, Zvereva and collaborators employed the same fluorescent tag in a fluorescence anisotropy immunoassay to monitor colchicine levels in biological fluids.⁷⁵

To improve the probe performance, Arnold *et al.* synthesized two BODIPY-colchicine conjugates (**1d–1e**) in 2008, which proved effective in both biochemical assays and live-cell imaging.⁷⁶ Other fluorophoric groups such as dansyl⁷⁷ or nitrobenzoxadiazole (NBD)⁷⁸ moieties have also been attached to colchicine (**1f–1g**) or its analogue colcemid (**1h**). It is worth mentioning that in the context of high-resolution microscopy, Zhang and co-workers developed a new colchicine photoswitchable fluorescent probe (**1i**) in 2015 by attaching a spiropyran derivative to the secondary amino group on ring B of colchicine.⁷⁹ They validated this probe for high-resolution microscopy to quantitatively monitor trace amounts of tubulin in cancer cells.

Recently, Jurásek and colleagues reported a novel, green-emitting BODIPY-colchicine (**1j**) with high potency in order to unveil the involvement of different cellular membrane-based compartments in the inhibition of microtubule cytoskeleton dynamics by colchicine.⁸⁰

Beyond colchicine-based fluorescent probes, Leynadier and co-workers developed in 1993 two pteridine analogues, (*S*)-(–) and

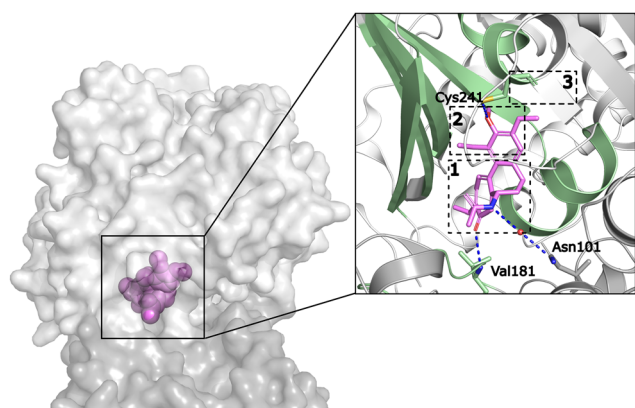


Fig. 2 Structure of colchicine-bound tubulin. (PDB: 5XIW). In the inset, the colchicine site is delineated by pale green ribbons and zones 1, 2 and 3 are boxed in dashed squares. Main interactions between colchicine and α -tubulin are shown by dashed lines. Highlighted are the hydrogen bonds with α -Val181 and β -Cys241 and a water-mediated hydrogen bond with α -Asn101. Oxygen and nitrogen atoms participating in the main interactions are shown in TV-red and blue, respectively.



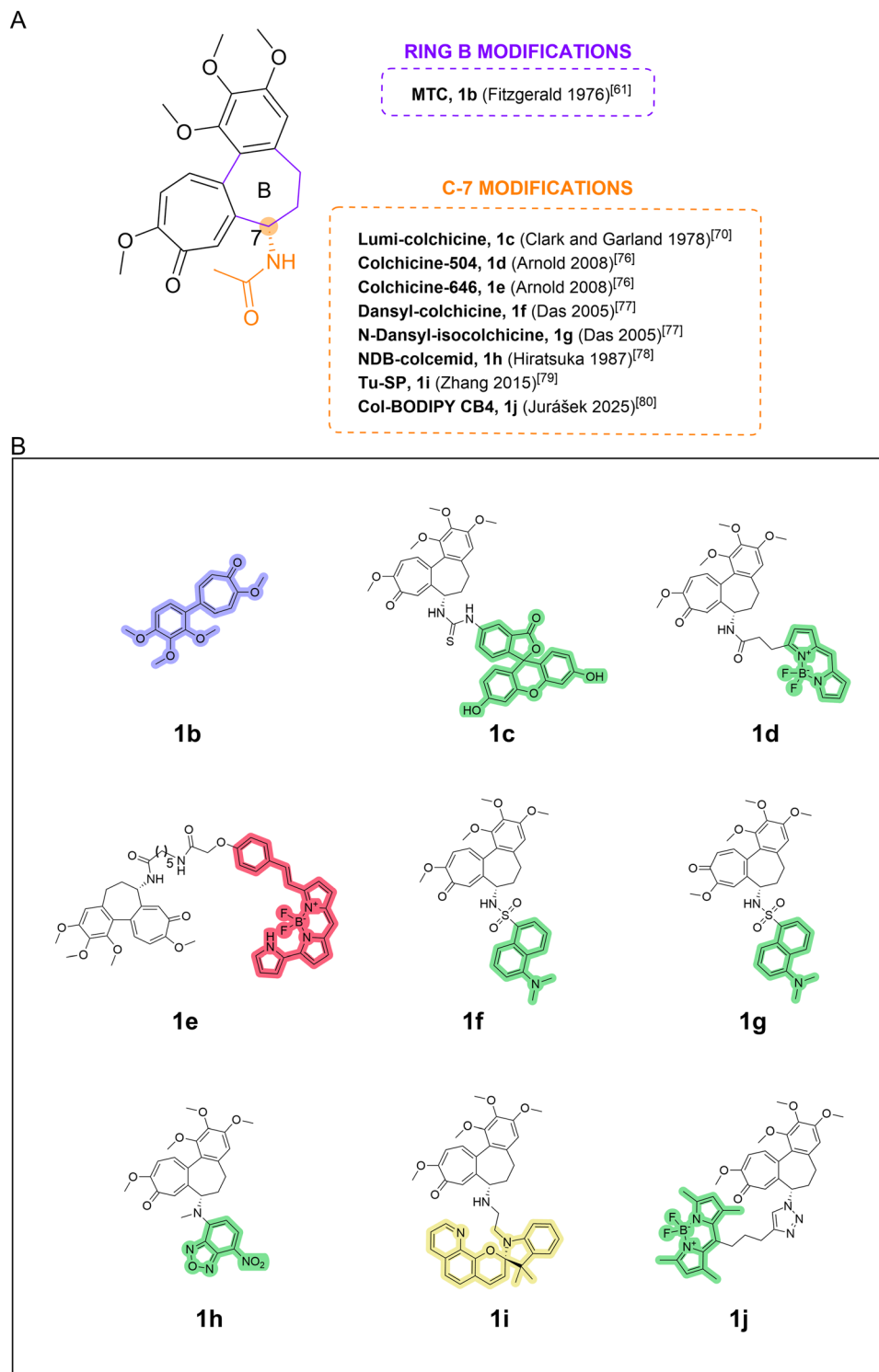


Fig. 3 Chemical structures of colchicine-based fluorescent probes. (A) Modification sites for fluorophore attachment on the colchicine core to yield biologically active fluorescent probes. (B) Chemical structures of colchicine-based fluorescent probes. The color of the highlighted regions represents the range of color emission of the corresponding fluorophore. RGB color codes assigned to represent each fluorescent emission range are as follows: UV (172,172,255); green (126,226,149); yellow (243,237,156); and far-red (251,98,124).

(*R*)-(+)-ethyl(5-amino-1,2-dihydro-2-methyl-3-phenylpyrido[3,4-*b*]-pyrazin-7-yl)carbamate (**2a** and **2b**, respectively) that bind to the colchicine site. These molecules fluoresce upon binding, with

affinities surpassing that of MTC ($4.1 \times 10^6 \text{ M}^{-1}$ for the (*S*)-isomer (**2a**) and $3.2 \times 10^6 \text{ M}^{-1}$ for the (*R*)-isomer (**2b**⁸¹)) (Fig. 4). The (*R*)-isomer in particular has become a widely used tool for



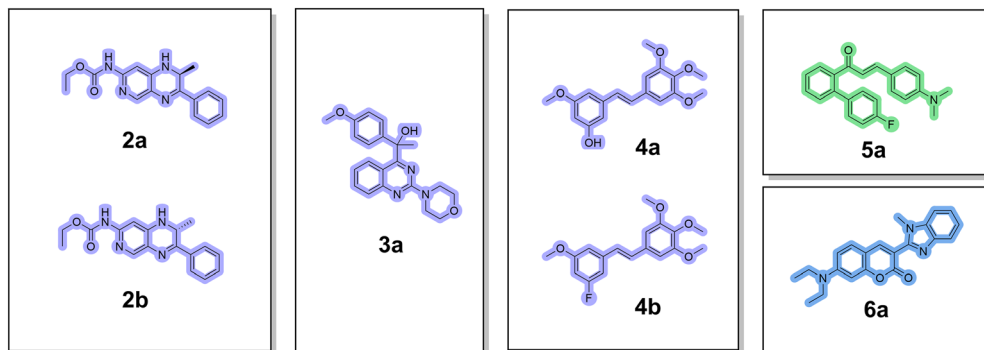


Fig. 4 Chemical structures of colchicine site fluorescent probes not based on the colchicine structure. Pteridines **2a** and **2b**, quinazoline **3a**, combreastatins **4a** and **4b**, chalcone OC9 **5a** and coumarin-30 **6a** are shown. RGB color codes assigned to represent each fluorescent emission range are as follows: UV (172,172,255); blue (119,173,237); and green (126,226,149).

identifying and characterizing high-affinity colchicine site binders.^{49,82–84} PM534, a novel colchicine site tubulin assembly inhibitor that occupies the entire binding pocket, was partially characterized using a competition assay with this fluorescent probe.⁵⁷ Additionally, Suzuki *et al.* developed a family of fluorescent quinazolines (**3a**) that bind the colchicine site both *in vitro* and in cellular systems.⁸⁵

Combreastatins—another group of colchicine site binders—exhibit potent antiproliferative activity. Among them, *Z*-isomers are more biologically active, but only the *E*-isomers exhibit fluorescence (**4a–4b**) (UV region). Leveraging this property, Bisby *et al.*⁸⁶ monitored compound uptake and intracellular distribution by means of fluorescence measurements (Fig. 4).

Additionally, in 2014⁸⁷ a series of *ortho*-aryl chalcones were characterized as a novel chemotype exhibiting tubulin-binding activity, with one derivative (OC9,⁸⁸ **5a**) displaying particularly potent antiproliferative capacity in both *in vitro* and *in vivo* experimental models. Furthermore, this compound exhibited fluorescence upon tubulin binding, thereby enabling its application as a fluorogenic probe for its spatial localization within the colchicine pocket in tubulin by fluorescent competitive assay. The authors employed this probe to efficiently characterize tubulin dynamics in different types of cancer cells.

Finally, a recent study⁸⁹ has identified coumarin-30 (**6a**), a laser dye, as a competitive ligand for the colchicine-binding site on tubulin. This compound exhibits substantially enhanced fluorescence properties, which make it a more versatile probe. This new ligand enabled the authors to identify and characterize a series of tubulin binders of the canonical colchicine-binding site and discover ligands that interact with other, less characterized binding pockets (*i.e.* “ $\beta\alpha I$ ” site).

Vinca domain

First isolated from the Madagascar periwinkle (*Catharanthus roseus*, previously *Vinca rosea*) in the mid-1950s by Cutts, Beer and Noble, vinblastine is one of the first anticancer agents used clinically.⁹⁰ The discovery of the compound traces back to the use of teas made from periwinkle leaves in folk medicine, which were supposed to have antidiabetic properties. In an

attempt to validate this assumption, Cutts, Beer and Noble administered extracts from *C. roseus* leaves to rats, but they did not observe any lowering in the blood glucose levels. Rather, the animals exhibited severe leukopenia and bone marrow suppression. Activity-guided fractionation of the crude extract eventually led to the isolation of a crystalline compound with chemotherapeutic potential, which was named vincalkebostine, now known as vinblastine. Vinblastine was approved for cancer treatment in 1961, while vincristine, another vinca alkaloid, was approved in 1963.⁹¹ Following these discoveries, second-generation vinca alkaloids were developed, including vindesine (only approved in some countries) and vinorelbine (reviewed in a previous study⁹²). These compounds act as microtubule destabilizers and bind to the vinca domain of tubulin, a site located at the interface between two $\alpha\beta$ -tubulin heterodimers. Compounds targeting the vinca domain inhibit microtubule assembly by acting as structural wedges at the longitudinal interface between tubulin dimers (reviewed in previous studies^{3,93}).

As is the case for most of the tubulin-binding sites for low-molecular weight ligands, the vinca domain is structurally promiscuous. Besides vinca alkaloids, structurally unrelated compounds that target the vinca domain and exhibit similar microtubule-destabilizing activity have been identified. These include halichondrins,⁹⁴ auristatins and dolastatins.^{95,96} Among them, eribulin, a synthetic analogue of the marine natural product halichondrin B, was approved by the FDA in 2010 for the treatment of breast cancer (https://www.accessdata.fda.gov/scripts/cder/daf/index.cfm?event=overview_process&ApplNo=201532).

The vinca domain extends across regions from two adjacent tubulin heterodimers and includes overlapping regions targeted by both vinblastine and dolastatins.⁵ As part of microtubules, the domain faces the inner lumen and is involved in longitudinal protofilament contacts. The vinblastine-binding pocket is enclosed by the T7 loop, helix H10, and strand S9 of the α -subunit together with the C-terminal turn of helix H6 and the T5 and H6–H7 loops of β -tubulin from the adjacent heterodimer.^{97,98} In contrast, the dolastatin site extends towards helix H1 of the β -subunit and typically accommodates



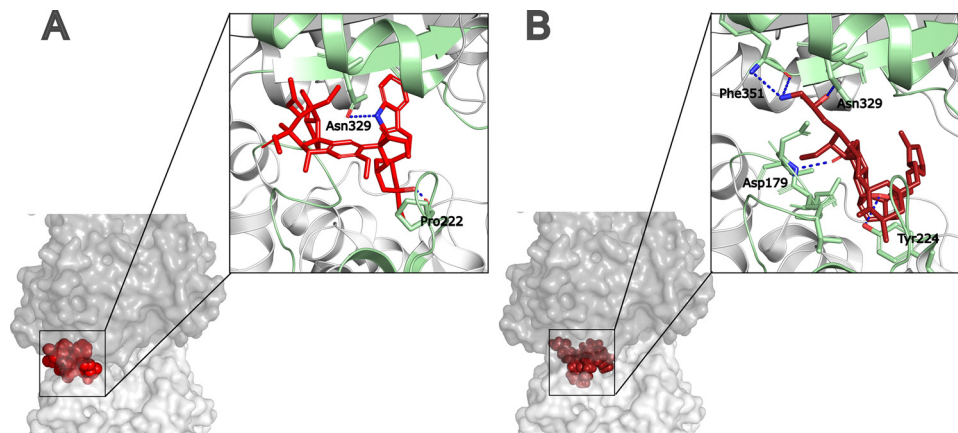


Fig. 5 Structures of vinca domain ligand-bound tubulin. Structure of (A) vinblastine-bound tubulin (PDB: 8CLE) and (B) eribulin-bound tubulin (PDB: 5JH7). In the inset, the vinca domain is delineated by pale green ribbons. Main interactions with α - and β -tubulin are shown as dashed lines. Highlighted are the (A) hydrogen bonds with β -Pro222 and α -Asn329 and (B) hydrogen bonds with β -Asp179, β -Tyr224, α -Asn329 and α -Phe351. Oxygen and nitrogen atoms participating in the main interactions are shown in TV-red and blue, respectively.

flexible peptides of three or more amino acid residues⁹⁹ (Fig. 5). The binding of vinca site ligands to this site stabilizes curved tubulin oligomers that are incompatible with the straight conformation required for microtubule polymerization. Additionally, some vinca site ligands disrupt GDP-GTP exchange in tubulin by interacting with β -Tyr224, a residue that stabilizes the guanine base interaction with the protein.^{99,100}

Several fluorescent probes have been developed based on vinblastine to study its binding site and interactions with tubulin (Fig. 6). In 1996, Rai and Wolff¹⁰¹ created the first fluorescent probe for the vinca site, a vinblastine-4'-anthranilate conjugate (**7a**), which fluoresces when binding to tubulin. Upon UV excitation at the absorption maximum of the anthranilate, the probe forms a covalent adduct with tubulin, enabling the localization of the vinca site, years before the first vinblastine-tubulin crystal structure was solved in 2005.⁹⁷

In 2002, Chatterjee *et al.*¹⁰² synthesized a coumarin-conjugated vinblastine (**7b**) to characterize its interaction with tubulin. Around the same time, BODIPY-vinblastine conjugates began to emerge. A commercially available version (originally developed by Molecular Probes, now Thermo-Fisher, but since discontinued (Cat. No. V12390)) became widely used (**7c**). Jiang *et al.*⁷² employed it in 1998 to assess whether a novel tubulin binder, 3-(iodoacetamido)benzoylurea, could compete for the vinca site. In 2002, Gruol, King, and Kuehne used the same probe to study P-glycoprotein interactions with vinca alkaloids,¹⁰³ and Lin and Chen later applied it to test the ligands of the pregnane X receptor (PXR), a key player in drug-drug interactions.¹⁰⁴ In 2014, they developed a high-affinity BODIPY-vindoline conjugate (**7d**) for further PXR-targeted screenings.¹⁰⁵

In the same year, Carney *et al.* synthesized a series of potent vinblastine analogues and characterized them using BODIPY-vinblastine (**7c**).¹⁰⁶ Several other studies have employed this BODIPY-vinblastine to some extent.^{107,108} Meimetis and colleagues¹⁰⁹ later designed additional BODIPY- and SiR-

vinblastine derivatives (**7e-7j**) to investigate their cellular uptake and distribution.

In 2014, the first fluorescent eribulin probe, which incorporates a BODIPY fluorophore attached to the C35-amino group, was reported by Laughney *et al.* (eribulin-BFL, **8a**).¹¹⁰ The probe retained biological activity, disrupted microtubules, and induced mitotic arrest in cancer cells. Nabekura *et al.*¹¹¹ used it to study the mechanisms of resistance to eribulin, demonstrating the direct link between eribulin resistance and MDR1 activity. Moreover, they showed that intravenous co-treatment with a nanoparticle-encapsulated MDR1 inhibitor enhanced eribulin-BFL accumulation in tumors. These results suggested that previous failures to overcome MDR1-mediated resistance were primarily due to the poor pharmacokinetic properties of MDR1 inhibitors, rather than the insufficient pharmacodynamic effect. In summary, the probe provided new insights into the mechanism of resistance against eribulin *in vitro* and *in vivo*, enabling the development of a strategy to overcome this problem.

More recently, a second fluorescent eribulin probe has been synthesized to explore drug resistance and microtubule dynamics (Fig. 7). Doodhi *et al.*¹¹² conjugated eribulin to Alexa Fluor 488 at the C35 position (**8b**), which based on the structural data of the tubulin-eribulin complex had been identified as an ideal site for fluorophore attachment. Using this probe, it was demonstrated that eribulin specifically binds to the plus end of microtubules, inhibiting growth at this end. Interestingly, while the minus ends were not blocked, they exhibited prolonged pauses. At nanomolar concentrations, single-molecule binding to protofilament ends was visualized, revealing how eribulin promotes microtubule catastrophe and disrupts dynamic instability.

Taxane binding site

The taxane binding site is an extended cavity situated on the luminal surface of β -tubulin, primarily constituted by



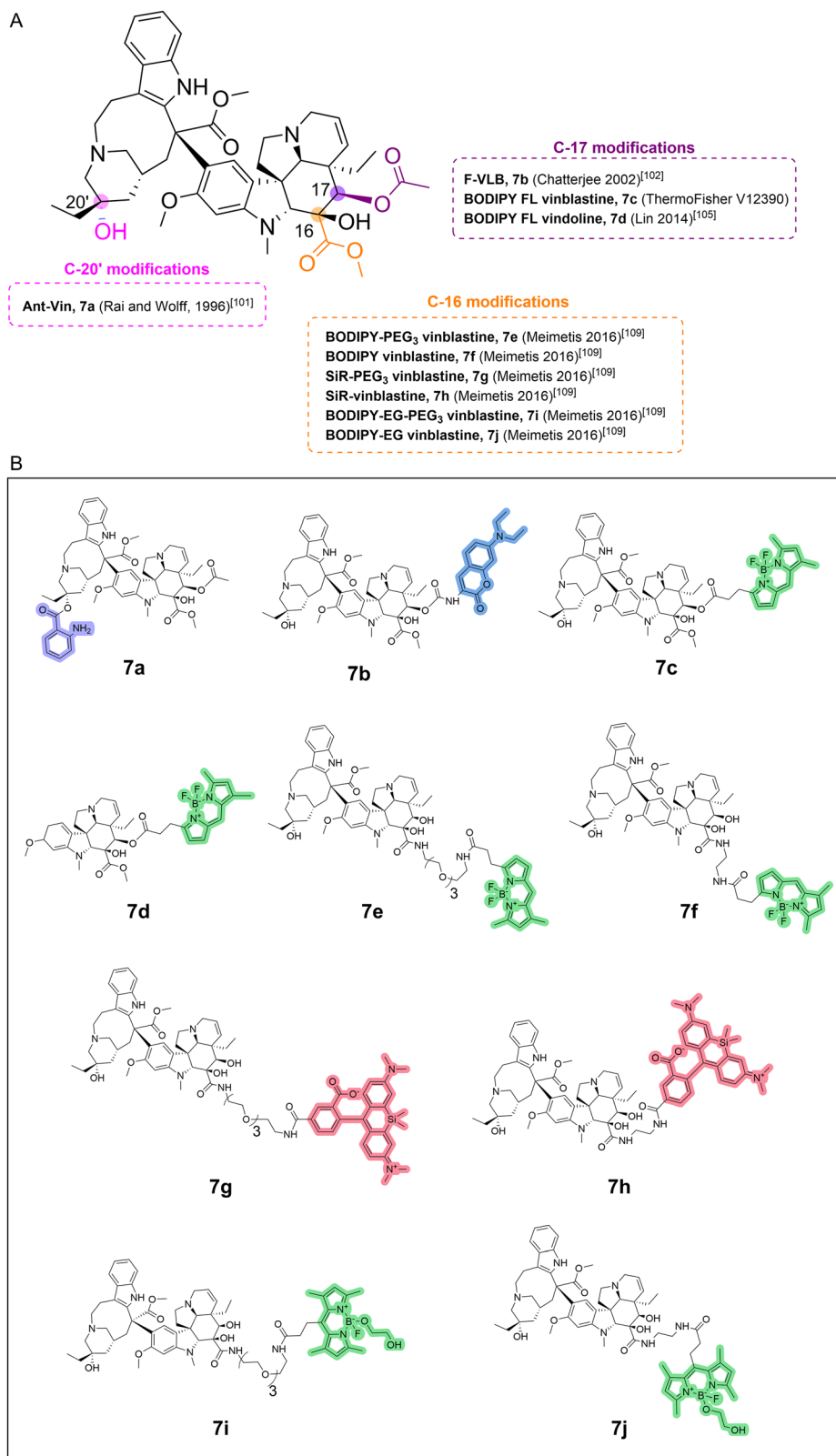


Fig. 6 Chemical structures of vinblastine-based fluorescent probes. (A) Modification sites for fluorophore attachment on the vinblastine core to yield biologically active fluorescent probes. (B) Chemical structures of these probes, with the color of the highlighted regions representing the range of color emissions of each attached fluorophore. RGB color codes assigned to represent each fluorescent emission range are as follows: UV (172,172,255); blue (119,173,237); green (126,226,149); and red (252,139,159).



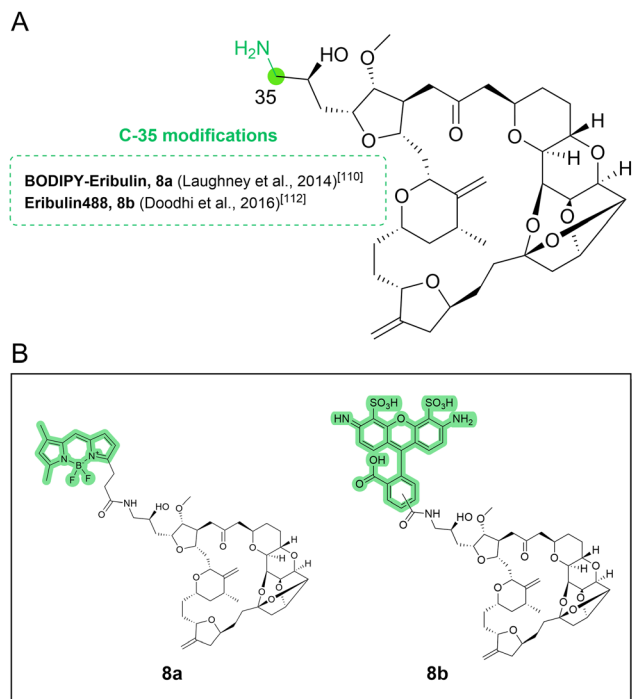


Fig. 7 Chemical structures of eribulin-based fluorescent probes. (A) Modification sites for fluorophore attachment on the eribulin core to yield biologically active fluorescent probes. (B) Chemical structures of these probes, with the color of the highlighted regions representing the range of color emission of each attached fluorophore. RGB color code assigned to represent green fluorescent emission range is (126,226,149).

hydrophobic residues from helices H1, H6, and H7, as well as the S9–S10 loop, the H6–H7 loop, and the M-loop (connecting

S7 and H9). A predominant effect of ligands binding to this site is the outward repositioning of the M-loop toward adjacent protofilaments, promoting favourable lateral contacts, and thus, stabilizing the microtubule and preventing its disassembly.^{7,113}

Several chemotypes have been shown to bind to this site in β -tubulin, with paclitaxel being the most extensively studied compound targeting this specific pocket. Since its clinical approval in 1992,¹¹⁴ the potent anticancer activity of paclitaxel has driven the development of a wide array of fluorescent taxane derivatives. These molecular tools have significantly advanced our understanding of taxane binding mechanisms and microtubule dynamics across various biological contexts. They have been instrumental in elucidating processes such as mitotic progression, cytoskeletal organization and cellular differentiation in diverse experimental models. Furthermore, these fluorescent probes have facilitated the identification and characterization of novel microtubule-targeting agents, contributing to the discovery of compounds with improved pharmacological profiles and reduced toxicity.

Similar to the vinca site, the taxane binding site is structurally promiscuous, and it accommodates the binding of several families of compounds, whose structures are not based on the taxane scaffold (Fig. 8). Notably, taccalonolides,¹¹⁵ zampanolides,¹¹⁶ epothilones¹¹⁷ and discodermolide¹¹⁸ have all demonstrated potent *in vitro* inhibition of cancer cell proliferation through this shared mechanism of microtubule stabilization.^{5,119} The discovery and characterization of these compounds have spurred the development of numerous structural analogues and derivatives,^{120–123} designed to improve the pharmacological properties and enhance the

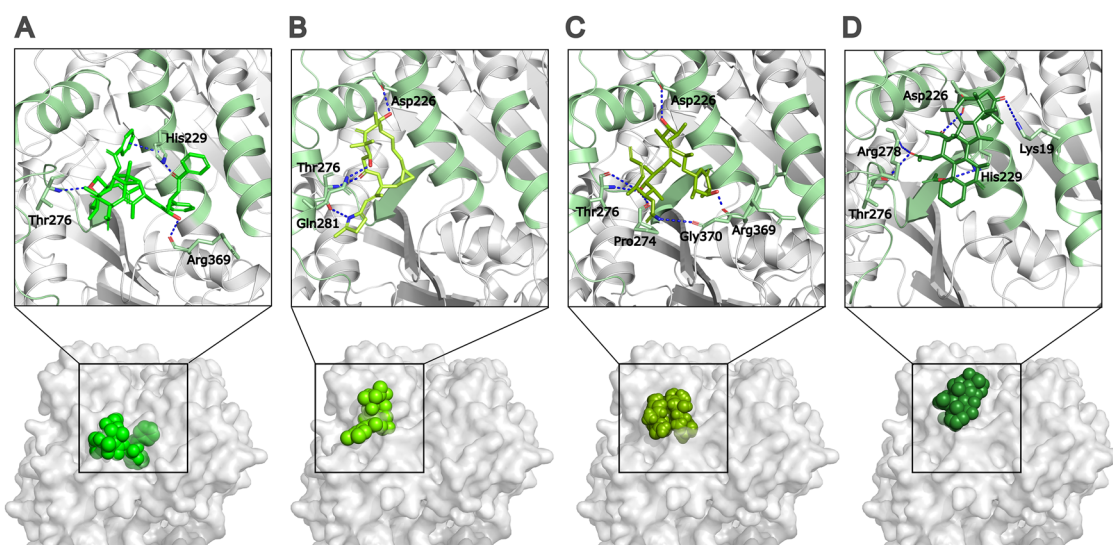


Fig. 8 Structures of taxane site ligand-bound tubulin. Structure of (A) paclitaxel-bound tubulin (PDB: 6WVR), (B) epothilone-bound tubulin (PDB: 4I50), (C) discodermolide-bound tubulin (PDB: 5LXT) and (D) taccalonolide-bound tubulin (PDB: 5EZY). In the inset, the taxane site is delineated by pale green ribbons. Main interactions with β -tubulin are shown as dashed lines. Highlighted are the (A) hydrogen bonds with β -Thr276, β -Arg369 and β -His229 and π -stacking with β -His229, (B) hydrogen bonds with β -Asp226, β -Thr276 and β -Gln281, (C) hydrogen bonds with β -Asp226, β -Thr276, β -Pro274 and β -Arg369, and (D) hydrogen bonds with β -Asp226, β -His229, β -Thr276 and β -Arg278 and a covalent bond with β -Asp226. Oxygen and nitrogen atoms participating in the main interactions are shown in red and blue, respectively.



synthetic accessibility. Fluorescent probes that are derived from taxanes, epothilones, discodermolide, or taccalonolides have been developed; however, probes based on the zampanolide scaffold are still missing at this point.

Taxanes. Paclitaxel was first discovered in the 1960s during a plant screening program conducted by the National Cancer Institute and the United States Department of Agriculture. It was isolated from the bark of the Pacific yew (*Taxus brevifolia*) and was approved by the FDA for the treatment of ovarian cancer in 1992. It was subsequently approved for breast cancer (1994) and non-small cell lung cancer (1999) (reviewed in a previous study¹¹⁴).

Although the detailed cellular and antitumoral mechanisms of action of paclitaxel remain subjects of ongoing investigations, it was demonstrated by Schiff *et al.* in 1979 that paclitaxel promotes microtubule polymerization.¹²⁴ Since then, paclitaxel has become one of the most successful drugs in the history of chemotherapy, and it has paved the way for the development of other taxanes as effective chemotherapeutic agents such as docetaxel, approved in 1996, and cabazitaxel approved in 2010 (<https://www.accessdata.fda.gov/scripts/cder/daf/index.cfm?event=overview.process&ApplNo=020449> and <https://www.accessdata.fda.gov/scripts/cder/daf/index.cfm?event=overview.process&ApplNo=201023>, respectively).

As one of the most widely used tubulin-targeting compounds, paclitaxel's binding mode has been extensively studied and exploited for the development of fluorescent derivatives. Investigations into the molecular mechanism of tubulin binding,^{125–127} and the structural environment in the binding site have inspired the synthesis of multiple fluorescent paclitaxel derivatives that retain biological activity (Fig. 9). Given paclitaxel's synthetic complexity, which is a common feature among MTAs, a compromise must be reached between synthetic feasibility and biological activity preservation.

Paclitaxel anchors in a deep hydrophobic cleft where it establishes hydrogen bonds and hydrophobic interactions, primarily involving the C3' side chain, C2, and C4 moieties. These interactions stabilize microtubules by reinforcing lateral protofilament association and inhibit depolymerization by precluding the straight-to-curved transition.¹²⁸

The total synthesis of paclitaxel became a hallmark challenge in organic chemistry, culminating in 1994 in the first successful total syntheses reported independently by the groups of Nicolaou and Holton, respectively.^{125,129} The pursuit of efficient synthetic routes remained ongoing in subsequent years, yielding 11 total syntheses reported to date, the most recent being published in 2021.¹³⁰ The current industrial production route of paclitaxel involves a semisynthetic approach that relies on 10-deacetyl baccatin III as the starting material; the latter is a natural product extracted from the leaves of the European yew (*Taxus baccata*).^{131,132}

Although the creation of high-affinity fluorescent probes does not require *de novo* total synthesis, the development of such derivatives needs meticulously designed synthetic strategies, including the use of appropriate protecting groups and optimized reaction conditions. In this section, we group

fluorescent derivatives according to the fluorophore attachment site on the taxane core.

C7 modifications

The C7 position emerged as the most useful site for chemical modification, yielding the highest number of biologically active, high-affinity fluorescent derivatives (Fig. 9). The C7 hydroxy group can be readily derivatized by means of esterification and is structurally positioned in such a way that its modification does not substantially impair the tubulin-binding affinity. The earliest example of a C7-modified fluorescent paclitaxel derivative was reported by Kingston *et al.* in 1991,¹³¹ who synthesized a dansyl-labeled paclitaxel (Dan-PTX, **9a**) using a β -alanyl linker to connect the fluorophore to the C7 hydroxy group. Although this compound exhibited low fluorescence, it marks a foundational step in the field.

The first major breakthrough came with the development of flutax-1 (**9b**), synthesized by Souto *et al.* in 1996.¹³³ This derivative incorporated carboxyfluorescein and was obtained by reacting the free amino group of 7-(β -alanyl)paclitaxel¹³⁴ with the N-hydroxysuccinimidyl ester of 4'-carboxyfluorescein. Flutax-1 was later employed by Evangelio *et al.* (1998)¹³⁵ in pioneering studies that demonstrated the utility of fluorescent taxanes for specific, efficient, and precise labeling of cellular microtubules.

In parallel, Evangelio *et al.* also evaluated other C7-modified derivatives with worse biochemical and biological properties and fluorescence intensity, including cutax,¹³³ 2'AcFlutax¹³⁶ and NBD derivatives of docetaxel modified at positions 3 and 7, which exhibited minimal biological activity.¹³⁷ A rhodamine-conjugated derivative named rotax (**9c**)¹³⁸ showed some potential in promoting microtubule assembly. However, its use in live, unfixed cells led to non-specific staining, including labeling of non-microtubular vesicles, highlighting the challenge of achieving specificity and retention of potent biological activity properties in probe design. As a result, the development of a C7-modified red-emitting paclitaxel probe remained an unmet need in the field for several years.

Further optimization of flutax-1 led to the synthesis of a difluoro-fluorescein-conjugated taxane derivative, flutax-2 (**9d**),¹³⁹ which exhibited enhanced photostability and slightly higher affinity than first-generation **9b**. Initial kinetic studies of flutax-1 and flutax-2 binding to microtubules revealed that the paclitaxel-binding site was readily accessible to these fluorescent taxanes, an unexpected finding given that the site is located within the microtubule lumen. These findings, supported by the use of both flutax-1 and flutax-2, prompted the development of an alternative model wherein paclitaxel accesses its luminal binding site through lateral diffusion across the microtubule lattice rather than through end-on diffusion from the microtubule tip.¹⁴⁰

Further evidence for this model came from the use of a more recent fluorescent derivative featuring a six-carbon linker between the amino group of 7-(β -alanyl)paclitaxel and the fluorophore (hexaflutax, **9e**),¹⁴¹ which provided robust support for the localization of these fluorescent taxoids at a



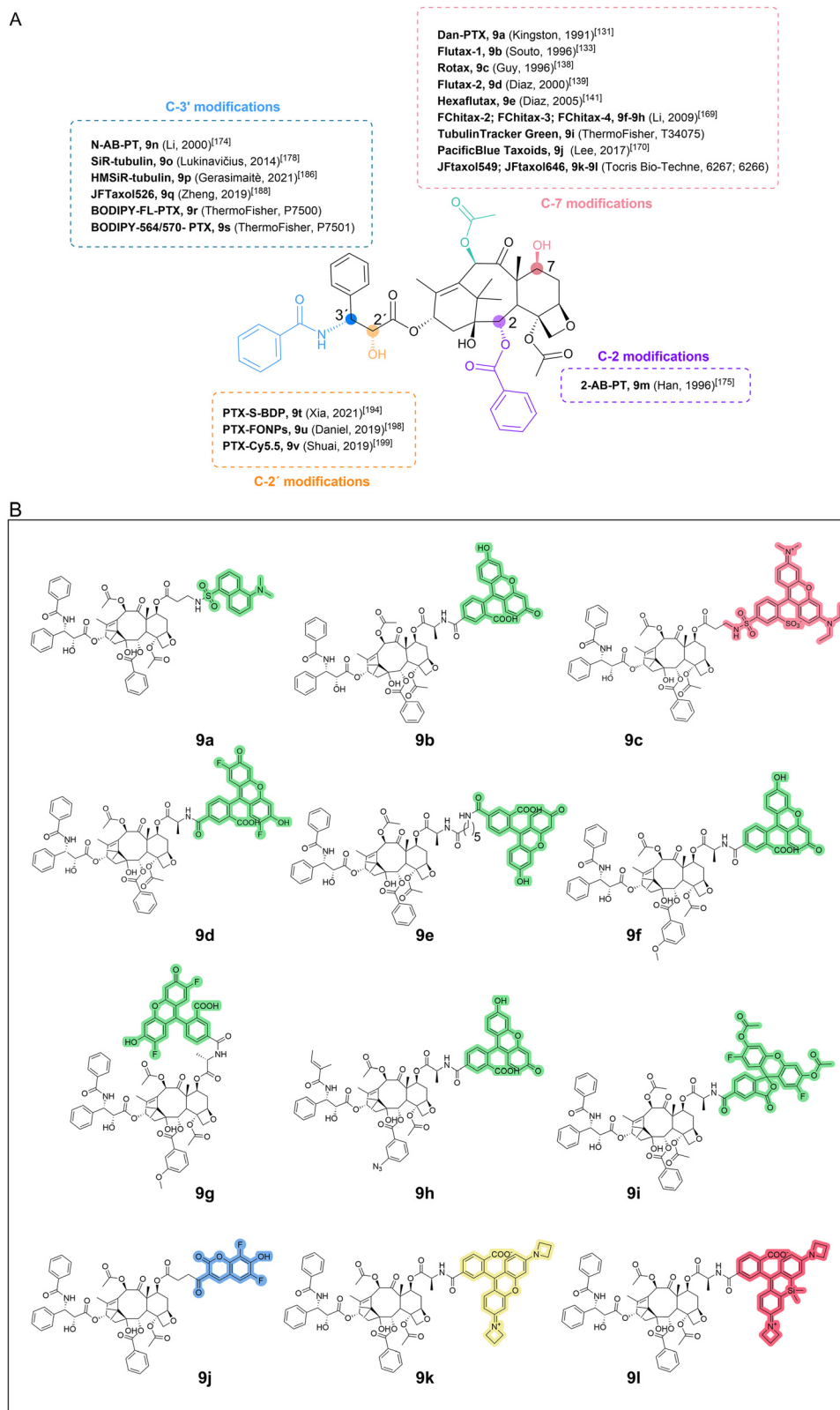


Fig. 9 Chemical structures of taxane-based fluorescent probes (I). (A) Modification sites for fluorophore attachment on the paclitaxel core to yield biologically active fluorescent probes. (B) Chemical structures of C7-modified probes, with the color of the highlighted regions representing the range of color emission of each attached fluorophore. RGB color codes assigned to represent each fluorescent emission range are as follows: blue (119,173,237); green (126,226,149); yellow (243,237,156); red (252,139,159); and far-red (251,98,124).



surface-accessible site that overlaps with the canonical paclitaxel binding pocket.^{142–144}

Among its many applications, flutax-2 has proven particularly useful in efficient identification of novel compounds targeting the paclitaxel binding site.^{145–147} When used with glutaraldehyde-stabilized microtubules,¹⁴⁸ flutax-2 enables robust competitive binding assays for evaluating MSAs.¹⁴⁹ Fluorescence anisotropy readouts from these assays allow the indirect determination of binding constants (K_b) for novel ligands with high accuracy and reproducibility.

In combination with sedimentation assays, flutax-2 has also facilitated the biochemical and kinetic characterization of ligand–tubulin and ligand–microtubule interactions for a wide array of compounds. These comprise cyclostreptin¹⁴² epothilones,^{150,151} dactyloides¹⁵² and zampanolides,^{62,122,147} and other taxane analogues developed by Matesanz and colleagues.^{153,154} Importantly, other MSAs such as peloruside and laulimalide did not displace flutax-2 from microtubules, thus revealing the existence of an alternative site within β -tubulin, now recognized as the laulimalide/peloruside binding site.^{155–157}

Since it was first introduced, flutax-2 has been widely adopted not only to investigate the cellular effects of paclitaxel but also to study the microtubule dynamics^{158,159} and the intracellular distribution of microtubules in different model systems.^{160,161} It has also enabled studies on cellular uptake, trafficking, and accumulation kinetics of taxanes,^{162,163} and has provided insights into microtubule-MAP interactions¹⁶⁴ and multidrug resistance (MDR) mechanisms.^{165–168}

However, accurate thermodynamic characterization using competitive binding assays with flutax-2 is limited to ligands whose affinities lie within three orders of magnitude of the probe. While flutax-2 can identify high-affinity binders, it cannot reliably determine their exact K_b values. Addressing this issue, Li *et al.*¹⁶⁹ developed a new series of high-affinity fluorescent probes, collectively named Fchitax. These derivatives incorporate fluorescein (**9f**) or oregon green (**9g**) fluorophores and were built upon paclitaxel scaffolds bearing methoxy substitutions at the meta-position in the phenyl ring of the C2-benzoyl moiety. In parallel, cephalomannine-based probes were synthesized by introducing an azide group at the same position (**9h**). These probes allow high-sensitivity detection at sub-nanomolar concentrations while improving the precision of affinity measurements for high-affinity taxane ligands.

Thermo-Fisher commercializes flutax-2 under the designation *Paclitaxel Oregon Green* (Cat. No. P22310), as well as a structurally related analogue featuring biacetylation of the phenolic hydroxyl groups within the fluorophore moiety, marketed as *Tubulin Tracker Green*, **9i** (Cat. No. T34075). This last derivative is an uncharged, non-fluorescent and more membrane-permeable compound compared to flutax-2. Upon cellular entry, the hydrolysis of ester bonds by endogenous nonspecific esterases removes the lipophilic acetyl groups, releasing the charged, green-fluorescent paclitaxel conjugate, i.e., flutax-2. Of note, the commercial kit of **9i** also includes

pluronic acid, to further enhance the cellular uptake of the probe.

In 2017, Lee and colleagues¹⁷⁰ expanded the fluorophore toolkit by attaching pacific blue (PB), a coumarin-based fluorophore, to the C7-hydroxy group of paclitaxel. The resulting PB-taxane probes emit at 447 nm, broadening the spectral range for multicolor imaging. Lee and coworkers systematically tested different linkers, glycine, β -alanine, and γ -aminobutyric acid (GABA), and found that the derivative with the shortest linker (PB-Gly-TXL, **9j**) conferred the highest affinity for microtubules.¹⁷¹ These PB-based probes were shown to be highly sensitive to cellular efflux mechanisms and required efflux pump inhibitors such as verapamil, which enhances intracellular retention, to achieve effective cytotoxicity in HeLa cells. Despite their size and spectral advantages,¹⁷² the cellular uptake of PB probes was less efficient than that of flutax-2, which demonstrated effective microtubule labeling without efflux modulation.

Lastly, C7-modified taxane probes incorporating Janelia Fluor (JF) dyes are now commercially available from Tocris Bio-Techne. These include JFtaxol549 (**9k**) (Cat. No. 6267) and JFtaxol646 (**9l**) (Cat. No. 6266), which emit in the yellow and far-red regions, respectively. Tocris also offers another taxane-binding probe, JFtaxol526, that shares similar optical properties. Notably, unlike the previous two conjugates, this compound is modified at the C3' position. The latter will be discussed in the next section.

C3', C2', C2 and C10 modifications

In addition to C7 labeling, several studies have explored fluorescent tagging of the paclitaxel molecule at alternative positions (Fig. 10), including C3' (Sengupta 1995,¹⁷³ Li 2000¹⁷⁴), C2 (Han 1996¹⁷⁵), or C10 (Baloglu 2001,¹⁷⁶ Sengupta 1997¹⁷⁷). In this context, it is worth mentioning the development of two minimal probes, 2-AB-PT¹⁷⁵ (**9m**) and N-AB-PT¹⁷⁴ (**9n**), containing an aniline group at C2 and C3, respectively. The studies of these probes paved the way for the development of SiR-tubulin (**9o**),¹⁷⁸ which enabled a significant advancement in microtubule imaging. This probe is based on docetaxel modified at the C3' position with silicon-rhodamine (SiR), a fluorophore in which the oxygen at position 10 of the xanthene ring system is replaced by silicon. The resulting Si-C bond imparts enhanced photostability and quantum yield, making SiR a highly attractive fluorophore for advanced imaging applications.^{179–181}

SiR-tubulin has been widely employed to investigate the microtubule structure and dynamics across a wide range of experimental models.^{182,183} Magliocca *et al.*,¹⁸⁴ for instance, used SiR-tubulin alongside SiR-actin¹⁷⁸ to study the interplay between actin and microtubule dynamics in induced pluripotent stem cells (iPSCs) and their neuronal derivatives. Their findings highlighted a reciprocal relationship between these cytoskeletal elements during differentiation.

More recently, Shen *et al.*¹⁸⁵ have utilized SiR-tubulin to analyze cytoskeletal remodeling in response to changes in osmolarity and molecular crowding. This work revealed that



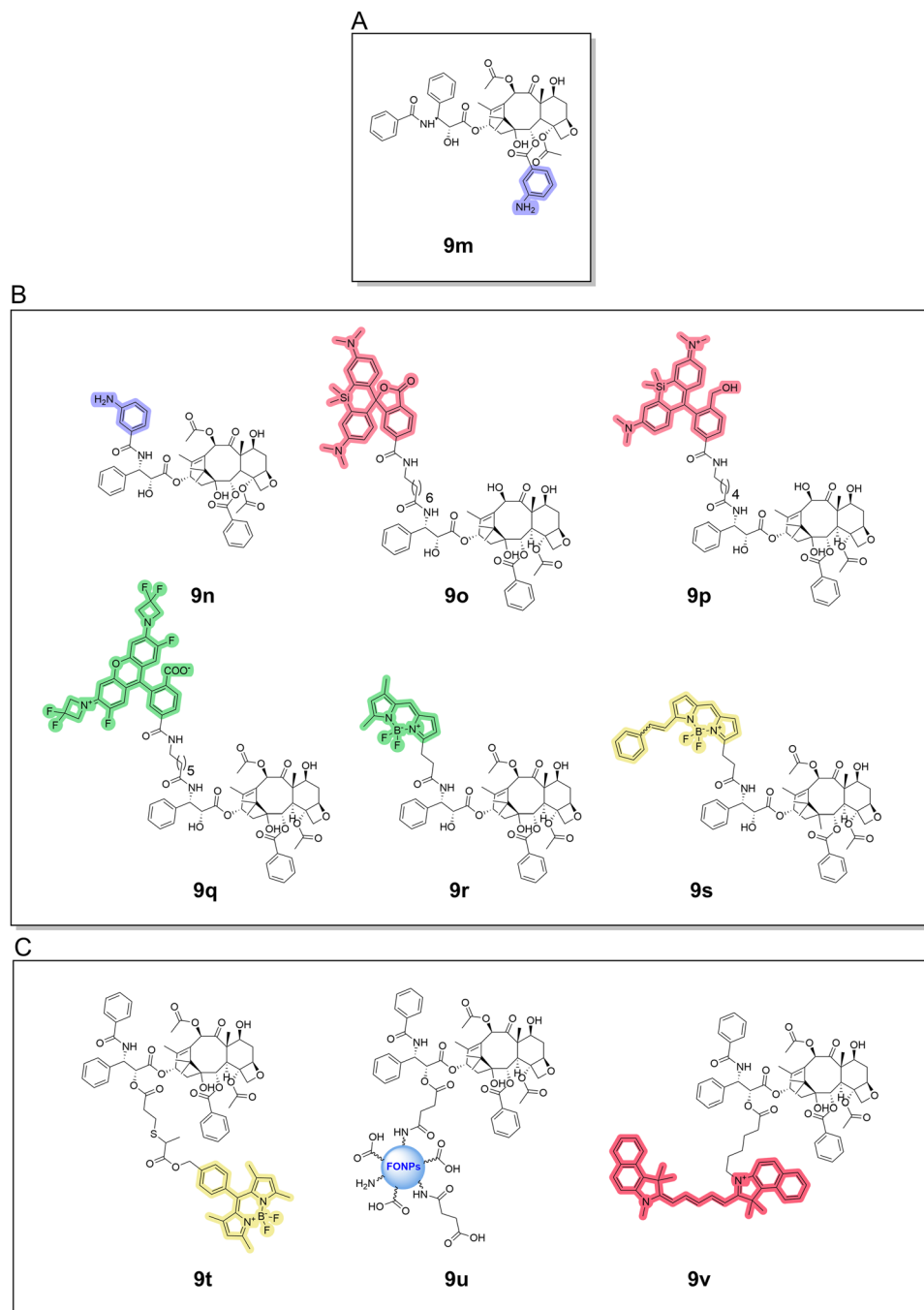


Fig. 10 Chemical structures of taxane-based fluorescent compounds (II). Chemical structures of (A) C3'-, (B) C2- and (C) C2'-modified probes, with the color of the highlighted regions representing the range of color emission of each attached fluorophore. RGB color codes assigned to represent each fluorescent emission range are as follows: UV (172,172,255); green (126,226,149); yellow (243,237,156); red (252,139,159); and far-red (251,98,124).

the cytoskeleton exhibits a highly dynamic capacity to respond to environmental stimuli through post-translational modifications such as acetylation and detyrosination, as well as through the regulated recruitment or release of microtubule-associated proteins, notably MAP7.

In 2021, a comparative study¹⁸⁶ focused on the coupling of various taxanes to hydroxymethyl silicon-rhodamine (HMSiR). The latter is a far-red dye known for its spontaneous blinking

properties *via* intramolecular spirocyclization,¹⁸⁷ making it well-suited for super-resolution techniques. The authors evaluated conjugates of different taxanes, including docetaxel, cabazitaxel, and larotaxel, ultimately identifying an optimal probe for nanoscopy applications. The most effective construct, cabazitaxel linked to the 6'-regioisomer of HMSiR *via* a six-carbon spacer (**9p**), enabled high-resolution visualization of microtubule ultrastructure, including the resolution of the



inner microtubule diameter, using techniques such as single-molecule localization microscopy (SMLM), stimulated emission depletion (STED), and MINFLUX nanoscopy.

In addition to SiR and HMSiR-based probes, Zheng *et al.*¹⁸⁸ developed a novel fluorogenic paclitaxel-site probe using Jane-lia Fluor 526 (JF526),^{189,190} yielding JFtaxol526 (**9q**). This probe is based on fluorophore attachment *via* C3' of the paclitaxel side chain, and it incorporates a green-emitting fluorophore exhibiting fluorogenic behavior analogous to SiR dyes *via* a lactone-zwitterion equilibrium. JF526 spontaneously transitions from a non-fluorescent spiroether form to a fluorescent zwitterionic state upon protonation at physiological pH. *In vitro* assays demonstrated that JFtaxol526 promotes tubulin polymerization, consistent with the mechanism of paclitaxel and its analogues. In live-cell experiments, the probe specifically stained microtubules in the presence of verapamil. However, detailed thermodynamic binding characterization was not reported by Zheng *et al.*¹⁸⁸

JFtaxol526 is now commercially available through Tocris Bio-Techne, along with JFtaxol549 and JFtaxol646, which emit in the yellow and far-red spectral ranges, respectively (*vide supra*).

Finally, Thermo-Fisher has also previously commercialized paclitaxel-BODIPY conjugates, including BODIPY-FL (**9r**) (green emission; Cat. No. P7500) and BODIPY 564/570 (**9s**) (orange emission; Cat. No. P7501); however, these products have recently been discontinued. Despite limited characterization in the literature, these probes, which are known to be substrates of P-glycoprotein,¹⁹¹ have contributed to our understanding of MSAs in cellular environments.¹⁹² For instance, a 2002 study employing BODIPY-FL-paclitaxel demonstrated that P-gp is a major barrier to paclitaxel transport across the blood-brain barrier.¹⁹³

Recent advancements in BODIPY-based probes have also facilitated new drug delivery strategies. Notably, Xia *et al.*¹⁹⁴ engineered a photoactivatable paclitaxel-BODIPY prodrug (PTX-S-BDP, **9t**) incorporating a redox-sensitive linker in C2' to enable fluorescence imaging-guided chemotherapy. Formulated as nanoparticles using pluronic F-127,¹⁹⁵ PTX-S-BDP exhibited improved aqueous solubility, high colloidal stability, and cytotoxicity comparable to unmodified paclitaxel. A key innovation of this system is the use of a sulfide [or thioether] linker sensitive to intracellular glutathione and hydrogen peroxide levels, facilitating selective activation in the tumor microenvironment.^{196,197}

Within the context of taxane-based nanoparticle systems, two additional formulations incorporating fluorescent paclitaxel derivatives have been previously developed. In 2019, Daniel *et al.* conjugated fluorescent organic nanoparticles, synthesized from citric acid and diethylenetriamine, to the 2'-hydroxy group of paclitaxel (**9u**).¹⁹⁸ The resulting water-soluble probe demonstrated preferential uptake by glioblastoma cells in two-photon fluorescence imaging assays. The second reported nanoparticle formulation consists of paclitaxel conjugated to Cy5.5 (**9v**), also *via* the C2' position.¹⁹⁹ This study, which primarily focused on evaluating the biodistribution of

paclitaxel, employed various nanoparticle formulations. Following intravenous administration in Bel-7402 tumor-bearing mice, the fluorescent conjugate enabled visualization of the tissue distribution of paclitaxel.

Fluorescent taxanes with unknown structures

In addition to the previously mentioned fluorescent taxanes whose structures are publicly accessible, several commercial entities have introduced fluorescent taxane-derived probes whose chemical structure is not available. For instance, Tubulin Tracker Deep Red (**9w**) (Invitrogen, now Thermo-Fisher) is a far-red-emitting derivative of docetaxel, but its exact chemical structure remains undisclosed. The lack of biochemical and biological characterization data (*e.g.*, binding affinity, polymerization activity, IC₅₀) limits a thorough understanding of the biological effects of this probe. Nevertheless, the manufacturer reports rapid and specific microtubule labeling in live cells within 30 minutes, with no measurable cytotoxicity observed after 24 hours of incubation at working concentrations. The commercial kit includes probenecid to block organic anion transporters and improve probe retention.

Biotium has also developed a suite of fluorescent paclitaxel-based probes under the ViaFluor brand (**9x**). These include ViaFluor 405 (blue emission; Cat. No. 70064), ViaFluor 488 (green; Cat. No. 70062), and ViaFluor 647 (far-red; Cat. No. 70063). Although their exact chemical structures remain undisclosed, these probes are described as paclitaxel-derived and reported to exert minimal perturbation of microtubule dynamics and cell division, presumably due to low microtubule affinity. However, quantitative data regarding binding affinities or other biochemical and biological parameters are not publicly available.

In 2020, Logan and McCartney²⁰⁰ conducted a comparative study using *Drosophila* oogenesis to assess the performance of three far-red microtubule probes: Tubulin Tracker Deep Red, ViaFluor 647, and SiR-Tubulin. All three probes successfully labeled microtubules, but Tubulin Tracker Deep Red exhibited the highest fluorescence intensity, best signal-to-noise ratio, and fastest cellular uptake. It also maintained effective labeling at lower concentrations that minimized the disruption of F-actin (probably because it did not require the use of verapamil). Interestingly, although verapamil improved probe retention, it also increased actin cytoskeleton defects, suggesting potential off-target effects.

Merck, in turn, offers a green-labeled taxane under the trade name BioTracker488 (**9y**) (Cat. No. SCT142). However, the chemical structure of this compound and its biochemical and biological characterization have not been disclosed.

Besides, Spirochrome offers a series of paclitaxel-derived fluorescent probes trademarked as SPY-tubulin (**9z**), including SPY-555 tubulin (Cat. No. SC203), SPY-650 tubulin (Cat. No. SC503), and SPY-700 tubulin (Cat. No. SC603). The product datasheets supplied by the manufacturer do not include any additional structural, biochemical or biological data. It is worth mentioning that SPY555-tubulin and SPY650-tubulin have been increasingly employed to visualize microtubule architecture



and dynamics in live-cell imaging. Each probe has been cited in over 30 studies, contributing to critical insights across a wide range of biological systems and cellular processes.^{201–204}

Despite the growing availability of these fluorescent tools, significant challenges remain in correlating the molecular effects of these probes on microtubules with their broader cellular impacts. In the absence of comprehensive characterization, it is difficult to interpret cellular phenotypes following probe incubation. A notable example is flutax-2, which induces the formation of microtubules with altered interdimer spacings and protofilament numbers compared to paclitaxel (Bonato *et al.* 2025, in press). Furthermore, cells treated with flutax-2 display distinct phenotypes, underscoring that observed effects may stem not only from the MTA itself but also from the attached fluorophore or linker moiety.

Finally, the widespread adoption of the commercial probes is limited by their considerable cost, which restricts accessibility for many laboratories.

Importantly, to conclude the section on fluorescent taxanes, we note that last year, Wen *et al.*²⁰⁵ reported a series of docetaxel-derived probes leveraging the concept of bioorthogonal chemistry. They developed a functionalized C3' derivative bearing an azide handle, enabling *in situ* post-fixation reaction with fluorophores containing strained alkynes such as DBCO. In combination with super-resolution microscopy of mitotic spindles, this strategy revealed dark spindle zones undetectable by immunolabelling and preserved fluorescence after fixation.

This approach holds considerable promise, with potential applications across various contexts and as inspiration for developing new probes from other scaffolds based on the same concept.

Epothilones. Epothilones are a class of natural macrocyclic lactones originally isolated from the myxobacterium *Sorangium cellulosum*.²⁰⁶ Among naturally occurring epothilones, the epoxide-containing epothilones A and B are the most abundant. The corresponding olefins, epothilones C and D, are still highly active, but they are approximately 10-fold less potent cell growth inhibitors than epothilones A and B, respectively (reviewed in a previous study²⁰⁷). Epothilones bind to the taxane site on β -tubulin and promote microtubule polymerization and stabilization. Their potent antiproliferative activity led to the development of ixabepilone, a semisynthetic lactam analogue of epothilone B that was approved in 2007 for the treatment of metastatic breast cancer (<https://www.accessdata.fda.gov/scripts/cder/daf/index.cfm?event=overview.process&ApplNo=022065>). Unlike paclitaxel, ixabepilone is a poor substrate of multidrug resistance (MDR) efflux pumps and is effective against several tubulin isotypes, including the taxane-resistant β III-tubulin isotype.²⁰⁸

Structurally, all epothilones share a conserved binding mode in which they form hydrogen-bonding networks with residues Thr276 and Gln281 of the M-loop. These interactions stabilize lateral contacts between adjacent β -tubulin monomers by engaging the M-loop of one monomer with the H1–S2 loop of the neighboring protofilament. In addition, they establish several hydrophobic contacts within the taxane pocket.^{113,209}

The first fluorescently labeled epothilone derivatives (Fig. 11) were reported by Ganesh *et al.* in 2003, who introduced a meta-*N,N*-dimethylaminobenzoic acid moiety at C26 of epothilone D (**10a**) or C26 of C21-hydroxy epothilone D (**10b**), respectively. While being substantially less active than the parent compound epothilone D, these probes retained sufficient biological activity to enable fluorescence-based mechanistic investigations.²¹⁰

A significant advance was made by Gertsch *et al.* in 2009, who synthesized a fluorescent azathilone probe, where an NBD fluorophore was attached to a backbone-modified epothilone core (12-aza-epothilone) (**10c**).²¹¹ In this scaffold, the epoxide oxygen in epothilone A is removed and C12 is replaced by nitrogen, thus creating a functional handle for the easy attachment of exocyclic substituent groups, including fluorophoric moieties. The corresponding NBD conjugate exhibits a binding constant (K_b) for microtubule binding of $2.2 \times 10^6 \text{ M}^{-1}$ and retains microtubule specificity, while the cytotoxicity of the compound was 40- to 100-fold lower than that of epothilone A. Cytotoxicity was slightly diminished (approximately 3-fold) in a cell line overexpressing P-glycoprotein, indicating moderate susceptibility to efflux-based resistance.

In 2020, Rai *et al.* reported an AlexaFluor 488-containing epothilone B-derived probe (**10d**),¹⁵⁹ originating from the group of Altmann at ETH Zurich.²¹² In this probe, the fluorophore-linker construct was attached to C6 of the epothilone macrocycle, which had been deduced to be a favorable site for the perturbation-free attachment of bulky groups based on SAR studies^{213,214} and structural data for tubulin-epothilone complexes.²⁰⁹ The assembly of this conjugate involved the synthesis of an epothilone B analogue incorporating a 5-hexynyl group at position C6 of the epothilone macrocycle instead of the natural methyl group and subsequent conjugation with the fluorophore *via* Huisgen cycloaddition. The Alexa-labeled compound showed a K_b value of $1.5 \times 10^8 \text{ M}^{-1}$ for microtubule binding. However, no data on live-cell staining or cytotoxicity have been reported for the compound. Using the same modified epothilone B analogue as a precursor, a fluorescein-containing probe (**10e**) was also prepared, which, however, exhibited approximately 300-fold lower affinity for microtubules than the Alexa Fluor 488 conjugate.²¹² The disparity in affinity has been attributed to the differences in the linker length between the triazole moiety and the fluorophore, although this hypothesis awaits experimental confirmation.

Discodermolide. Discodermolide is a complex polyketide first isolated in 1990 from the marine sponge *Discodermia dissoluta*.²¹⁵ It exhibits potent antiproliferative activity²¹⁶ and stabilizes microtubules through binding to the taxane site on β -tubulin.²¹⁷ Due to its scarcity and the difficulty of sourcing the compound from natural specimens, numerous synthetic efforts have been undertaken to achieve its total synthesis.^{118,218–222}

The first high-resolution structural insight into discodermolide–tubulin interactions was provided by Prota *et al.* in 2017.²²³ Their study revealed that discodermolide binds deeply within the taxane pocket of β -tubulin, forming hydrogen bonds with residues Asp226, Ser232, Pro274, Thr276, and Arg369.



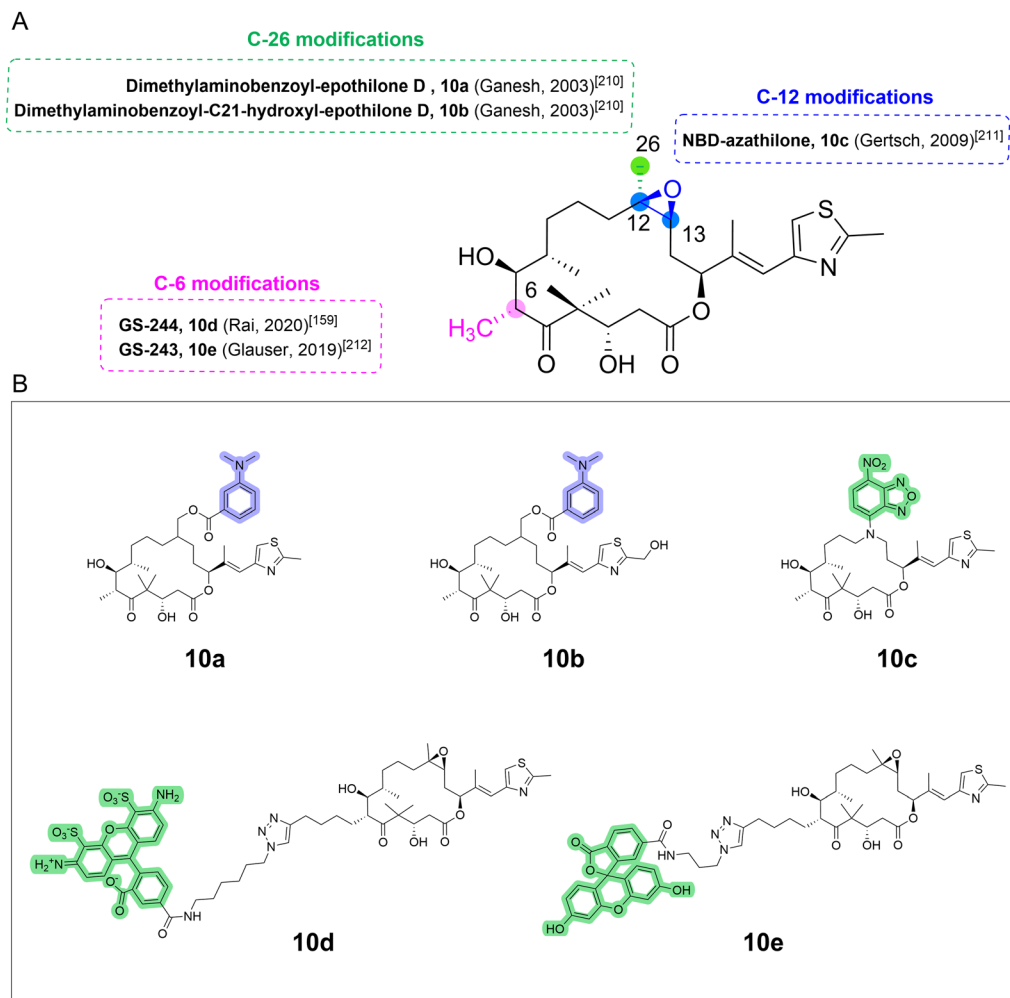


Fig. 11 Chemical structures of epothilone B-based fluorescent probes. (A) Modification sites for fluorophore attachment on the epothilone B core to yield biologically active fluorescent probes. (B) Chemical structures of these probes, with the color of the highlighted regions representing the range of color emission of each attached fluorophore. RGB color codes assigned to represent each fluorescent emission range are as follows: UV (172,172,255) and green (126,226,149).

Additional interactions involve the diene moiety engaging with Thr276 and a guanidyl group interacting with Arg278. These contacts contribute to the stabilization of the N-terminal segment of the M-loop by forming an α -helical conformation, a critical region for lateral protofilament contacts in microtubules. Moreover, van der Waals and hydrophobic interactions further anchor the compound within the binding pocket.

The first fluorescently labeled discodermolide derivatives (Fig. 12) were reported in 2005 by Smith *et al.* (**11a–11b**) alongside a series of five photoaffinity probes.²²⁴ The authors utilized a highly advanced intermediate from the total synthesis to generate two distinct fluorescent probes. One probe was obtained *via* chemoselective esterification of the primary hydroxy group of a partially protected discodermolide analogue with a hexanoic acid derivative bearing a dansyl fluorophore. The resulting construct retained both tubulin-polymerizing activity and cytotoxicity comparable to natural discodermolide. The second probe was obtained by the conjugation of N-6-aminohexyl dansylsulfonamide to the C19 position *via* a

carbamate linkage, producing a derivative with slightly diminished tubulin-polymerizing activity but enhanced cytotoxicity.

Subsequently, in 2011, a novel fluorescent derivative of a chemically simplified discodermolide analogue incorporating a dimethylaminobenzoyl (DMAB) moiety was developed (**11c**).²²⁵ This compound was designed to probe the microenvironment of the discodermolide binding site on tubulin and has potential utility in competitive binding assays for ligands with low affinity.

Taccalonolides. Taccalonolides are a family of microtubule-stabilizing compounds isolated from various *Tacca* species.^{226,227} Despite their ability to induce microtubule alterations reminiscent of paclitaxel's effects in live-cell systems at relatively high concentrations (5–10 μ M),¹¹⁵ their ability to affect microtubule polymerization *in vitro* has been found to be negligible.²²⁸ In 2017, Wang *et al.* solved the crystal structure of taccalonolide AJ bound to tubulin, which showed that the compound formed a covalent bond with tubulin by reaction between its epoxide group and β -Asp226.²²⁹



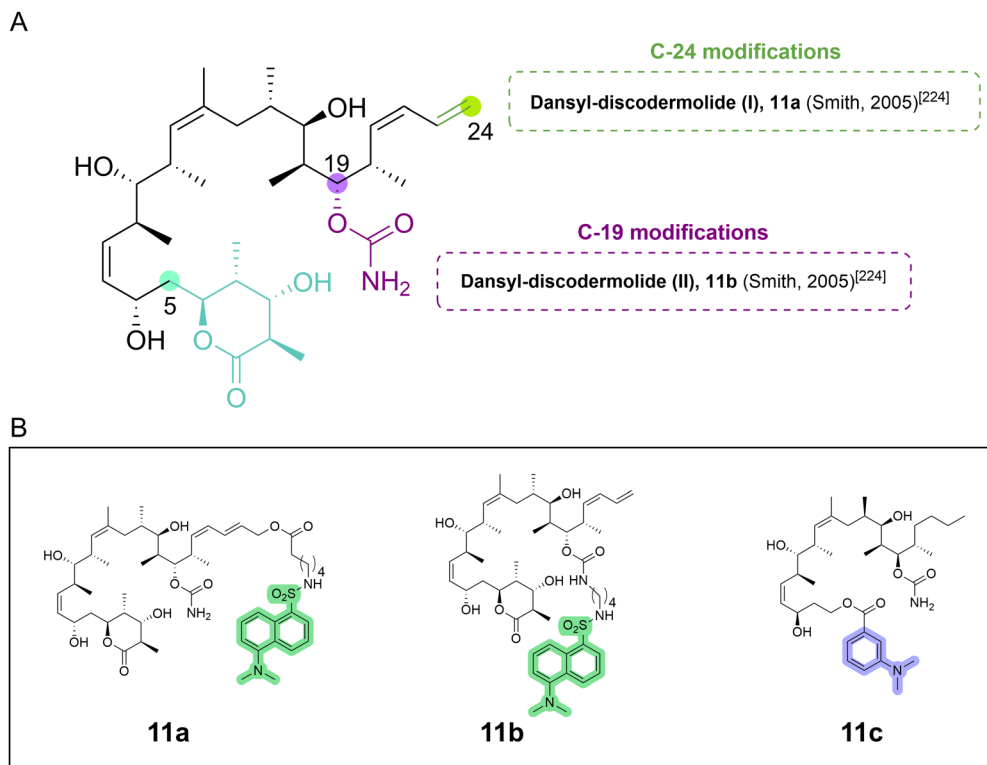


Fig. 12 Chemical structures of discodermolide-based fluorescent probes. (A) Modification sites for fluorophore attachment on the discodermolide core to yield biologically active fluorescent probes. (B) Chemical structures of these probes, with the color of the highlighted regions representing the range of color emission of each attached fluorophore. RGB color codes assigned to represent each fluorescent emission range are as follows: UV (172,172,255) and green (126,226,149).

Taccalonolides do not significantly enhance tubulin assembly and show poor competition with flutax-2 for microtubule binding.^{149,229} Nevertheless, they have attracted substantial interest in oncology due to their ability to circumvent resistance mechanisms commonly observed with taxane-based therapies. Notably, taccalonolides A, B, E, and N have been demonstrated in both *in vitro* and animal models to overcome resistance associated with P-glycoprotein overexpression, multidrug resistance-associated Protein 7, and β III-tubulin.^{230,231} A striking feature of these compounds is their *in vivo* potency, which surpasses expectations based on their *in vitro* activity.

In 2019, Du and colleagues identified C6 as an optimal position for fluorophore conjugation and they synthesized a taccalonolide–fluorescein conjugate (**12a**) through site-specific modification at this carbon.²³² Although this probe exhibited diminished cytotoxic potency in cellular assays, it retained comparable microtubule-binding capacity to the parental compound, laying the foundation for the development of future fluorescent taccalonolide probes.

Further work by the same group led to the synthesis of additional fluorescent derivatives that preserved both biochemical activity and cellular potency (Fig. 13).²³³ However, the applicability of these probes in live-cell imaging is limited by their irreversible binding to tubulin. Among the newly developed derivatives, Flu-Tacca-7 (**12b**), a C6-conjugated compound derived from taccalonolide AJ incorporating a

bis-pivaloyl-protected carboxyfluorescein, showed the most promising labeling properties. Flu-Tacca-7 confirmed β -tubulin as the primary molecular target of taccalonolide AJ and demonstrated robust labeling performance, particularly in contexts where non-covalent probes are ineffective, such as in drug-resistant cell lines expressing efflux transporters.

Maytansine site

Since its discovery in 1972,²³⁴ maytansine and related compounds have attracted attention as potent antiproliferative agents due to their unique mechanism of action on microtubules. However, only two maytansinoid-based compounds have been approved to date for clinical applications, both of which are antibody–drug conjugates (ADCs): trastuzumab emtansine (commercialized as KadcylaTM) consists of a maytansinoid linked to the monoclonal antibody trastuzumab, while mirvetuximab soravtansine-gynx (commercialized as ElahereTM) is a first-in-class antibody–maytansine conjugate targeting the folate receptor α (FR α).²³⁵ These ADCs have been approved for clinical use in HER2-positive metastatic breast cancer²³⁶ and FR α -positive, platinum-resistant ovarian cancer,²³⁷ respectively. Research continues to discover novel ligands targeting this binding site for potential clinical applications.

Maytansine-site ligands can bind to both curved and straight tubulin conformations and modulate microtubule dynamics through two distinct mechanisms. At high



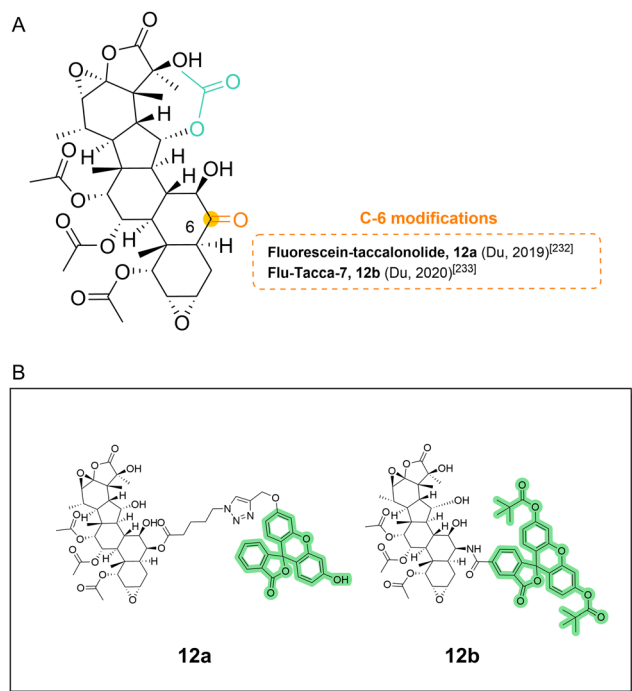


Fig. 13 Chemical structures of taccalonolide AJ-based fluorescent probes. (A) Modification sites for fluorophore attachment on the taccalonolide AJ core to yield biologically active fluorescent probes. (B) Chemical structures of these probes, with the color of the highlighted regions representing the range of color emission of each attached fluorophore. RGB color code assigned to represent green fluorescent emission range is (126,226,149).

concentrations, they sequester free tubulin into assembly-incompetent complexes. At sub-stoichiometric concentrations, they inhibit microtubule polymerization by sterically blocking the longitudinal addition of tubulin subunits at microtubule plus ends, thereby impairing polymer growth.²³⁸

Structural studies have clarified previous controversies regarding the mutually exclusive tubulin binding of maytansine and vinca domain ligands. It was unclear whether this inhibition was competitive or non-competitive. Crystallographic analyses by Prota and colleagues in 2014 resolved this paradox by demonstrating that maytansinoid binding hinders the complete assembly of the vinca domain, while the formation of the full vinca domain interferes with access to the maytansine site, indicating a mutually exclusive but non-competitive binding relationship.²³⁹ The maytansine binding pocket (Fig. 14) is located exclusively within β -tubulin and is formed by hydrophobic and polar residues from helices H3, H5, H11 and H11', and loops T3 and T11-11'.

The first fluorescent probe targeting this site (Fig. 15) was developed in 1993 from the 20-demethoxy-20-hydroxy derivative of ansamitocin P-3 (PDM-3).²⁴⁰ The conjugation of the phenolic hydroxy group with a dansyl derivative *via* a 3-carbon spacer yielded a fluorescent compound (**13a**) used in competitive assays to validate its binding specificity.

In 2015, BODIPY-labeled maytansinoids were introduced to facilitate the determination of drug-to-antibody ratios in

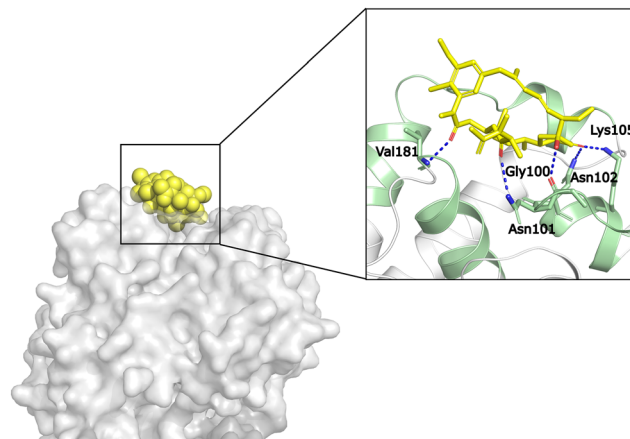


Fig. 14 Structure of maytansine-bound tubulin. (PDB: 4TV8). In the inset, the maytansine site is delineated by pale green ribbons. Main interactions with α -tubulin are shown as dashed lines. Highlighted are the hydrogen bonds with β -Gly100, β -Asn101, β -Asn102, β -Lys105, and β -Val181. Oxygen and nitrogen atoms participating in the main interactions are shown in TV-red and blue, respectively.

ADCs.²⁴¹ Two widely studied maytansinoids, namely DM1 and DM4, were conjugated through their thiol group at the terminus of the C3 side chain (**13b-13e**). One derivative displayed unexpectedly low fluorescence, probably due to photoinduced electron transfer quenching between the maytansinoid core and BODIPY.

A more functionally robust fluorescent probe was developed by Menchon *et al.* in 2018,²⁴² who synthesized a fluorescein-labeled maytansinoid derivative *via* the conjugation of the fluorescent moiety to an ester of maytansinoid with hept-6-ynoic acid by means of a click reaction with an azide-functionalized fluorescein derivative (**13f**). This probe demonstrated high affinity for tubulin ($K_b = 1.5 \times 10^8 \text{ M}^{-1}$) and faithfully mimicked the binding profile of maytansine. It was subsequently used to characterize the binding of compounds with unclear interactions, including spongistatin, disorazole Z, and PM060184.²⁴² Owing to its favorable properties, this probe has since been adopted in multiple studies,^{243,244} including the work by Bold *et al.*,²⁴⁵ who applied it to characterize a structurally simplified disorazole Z analogue.

Future perspectives. Developing biologically active probes derived from MTAs is a challenging task. Thus far, fluorescent probes have been developed for only 4 of the 8 druggable sites found in tubulin. Why are such molecular tools for the remaining four sites still lacking? While the gatorbulin and todalam sites have been discovered only recently, molecular probes are also absent for the laulimalide/peloruside site and the pirone-tin site, both of which have been known for much longer.

The laulimalide/peloruside site deserves special attention. The discovery of laulimalide and isolaulimalide (also named fijanolid B and A, respectively), independently reported by Corley *et al.*²⁴⁶ and Quiñoà *et al.*²⁴⁷ in 1988, introduced a new class of marine-derived macrolides with cytotoxic activity against tumor-derived KB cell lines, with laulimalide being the



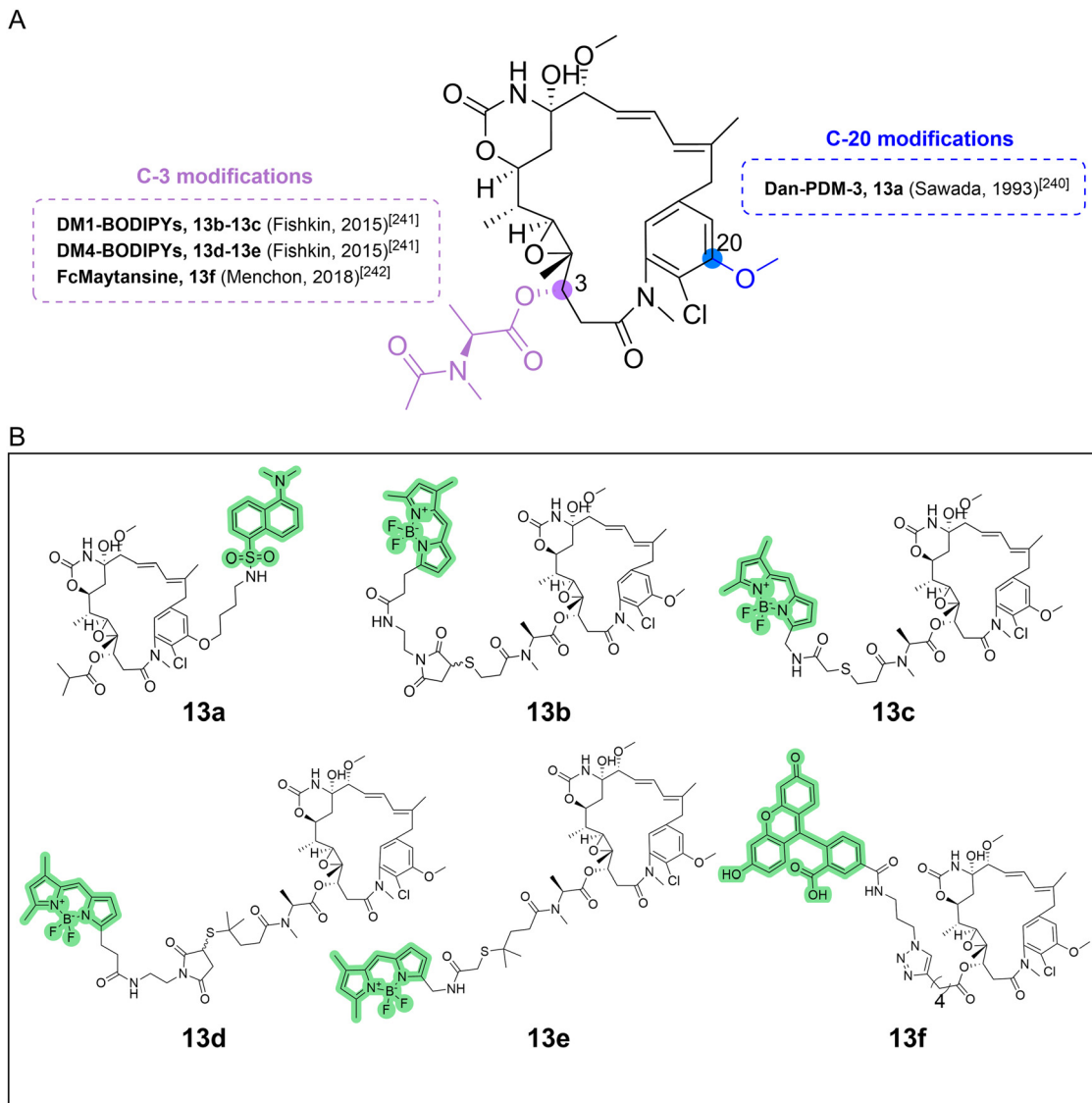


Fig. 15 Chemical structures of maytansine-based fluorescent probes. (A) Modification sites for fluorophore attachment on the maytansine core to yield biologically active fluorescent probes. (B) Chemical structures of these probes, with the color of the highlighted regions representing the range of color emission of each attached fluorophore. RGB color code assigned to represent green fluorescent emission range is (126,226,149).

more potent congener. Subsequent studies by Mooberry *et al.* in 1999 demonstrated that laulimalide stabilizes microtubules in cells and is a poor substrate of P-glycoprotein efflux pump.²⁴⁸

The marine macrolide peloruside A was first isolated from *Mycale* sp. by Northcote and co-workers²⁴⁹ in 2000. The same group subsequently demonstrated in 2002 that the compound possessed microtubule-stabilizing activity.²⁵⁰

By the early 2000s, laulimalide¹⁵⁵ and peloruside A¹⁵⁷ were hypothesized to bind to a shared site on microtubules that was distinct from the taxane pocket, based on competitive displacement assays. This hypothesis was later supported through multiple additional studies.²⁵¹⁻²⁵⁵ In 2014, Prota *et al.* solved the crystal structures of tubulin in complex with both compounds, confirming the existence of a novel outer surface binding pocket that is spatially distinct from the lumen-facing

taxane site. This configuration allows for the simultaneous binding of ligands at both sites without steric conflict.¹⁵⁶

Recently, Estévez-Gallego *et al.* described microtubule structural signals arising from the binding of laulimalide and peloruside.²⁵⁶ While the binding of taxanes leads to a longitudinal expansion of the tubulin dimer, laulimalide/peloruside is neutral with respect to dimer expansion/compression. In addition, it was shown that the structural effects of pelophen B,²⁵⁷ a synthetic analogue of peloruside A, do not even affect the lateral parameters of the microtubule lattice, thus extending the spectrum of compounds with potential clinical utility.

Notwithstanding the promising pharmacological relevance of laulimalide/peloruside, several challenges hinder probe development:

- The difficulty of chemical derivatization and limited availability of natural laulimalide and peloruside A.



- The synthetic complexity of *de novo* total synthesis of both parent compounds and suitably modified analogues.^{258–260}
- The tight steric environment surrounding the binding pocket, which complicates fluorophore incorporation.
- The limited accessibility of functional groups that are not directly involved in tubulin interactions, making derivatization chemically restrictive.

The pironetin site likewise remains unaddressed in the development of fluorescent probes. Pironetin was first isolated in 1994 by Kobayashi *et al.* from a *Streptomyces* sp. culture.^{261–263} While the compound was initially characterized as a plant growth regulator, subsequent studies by Osada and co-workers revealed its antiproliferative properties and its ability to induce cell cycle arrest.²⁶⁴ In 1999, Osada's group demonstrated that pironetin suppresses microtubule assembly.²⁶⁵ This compound covalently binds to a pocket in α -tubulin and destabilizes microtubules *via* two proposed mechanisms: (i) at high concentrations, pironetin forms assembly-incompetent tubulin-ligand complexes with free tubulin dimers and (ii) at low concentrations, it binds to the minus end of microtubules, where α -tubulin subunits are exposed, blocking further dimer addition. The development of probes targeting this site remains a compelling goal to advance the structural and pharmacological characterization of this unique binding pocket. The reason may lie in the site's location *per se*, which is set deeply embedded in the tubulin structure. This positioning limits both the diversity of molecular scaffolds reported to bind to this site and the possible chemical modifications of the pironetin scaffold that preserve its activity.

As pointed out above, compared to the laulimalide/peloruside and pironetin sites, the gatorbulin and todalam sites have been discovered more recently. This may partly explain why no fluorescent probes have been developed for these sites yet.

The gatorbulin site is named after the cyclic depsipeptide gatorbulin, which was isolated from the marine cyanobacterium *Lyngbya cf. confervoides* in 2021 by Mathew *et al.*²⁶⁶ The compound exhibits potent antiproliferative activity across several cancer cell lines by inhibiting microtubule dynamics. While the MTC probe (*vide supra*) has been employed to assess the affinity of gatorbulin and related analogues for tubulin,²⁶⁷ no site-specific fluorescent probe has yet been developed for its binding pocket. Although the gatorbulin site structurally overlaps with the colchicine site, the two sites are functionally divergent. Thus, the rational design of fluorescent ligands specific for the gatorbulin site would not only refine its biochemical characterization, but also allow spatiotemporal drug tracking in live-cell systems.

The only notable molecule reported so far to interact with the todalam site in α -tubulin is todalam itself, as discovered in 2022 by Mühlethaler *et al.* through a systematic approach combining high-throughput fragment-based crystallographic screening, iterative structure-based design, and extensive structure-activity relationship (SAR) analyses.²⁶⁸ Todalam has strong microtubule-depolymerizing activity, induces G2/M arrest in cells, and promotes apoptosis. This molecule acts

through a dual mechanism: being a dimer hijacker and functioning as a wedge, thus blocking the curved-to-straight transition and, hence, microtubule dynamics. In principle, the distinct location and mechanism of todalam offer a promising foundation for the rational development of site-specific imaging agents, which could provide novel tools for investigating microtubule dynamics and drug-target engagement in real time.

As this review demonstrates, a multitude of fluorescent probes binding to tubulin/microtubules have been developed over the years, which have enabled significant advances in our understanding of tubulin biochemistry and biology. The constant improvement of these probes required continuous optimization of both the linker moiety and the fluorophore. Early fluorophores, which incorporated simple polyaromatic systems or aromatic rings bearing electron-donating substituents (such as aniline), lacked the photophysical properties required for cytoskeletal visualization in advanced imaging applications. The development of probes incorporating more sophisticated fluorophores has now enabled the visualization of the cytoskeleton using techniques such as TIRF and high-resolution microscopy, which have revolutionized the microtubule field. In this context, numerous studies have employed diverse fluorescent probes to investigate the structural and dynamic properties of microtubules in the presence of various MTAs^{112,159,269} and associated proteins.²⁷⁰ However, it must still be kept in mind that modifying a parental MTA by the incorporation of a fluorophore can produce relevant unexpected effects, even if binding affinity and mode of interaction are maintained. An illustrative example is the case of flutax-2, which exhibits distinct effects on microtubules from the parent drug paclitaxel. Among other differences, flutax-2-assembled microtubules have a different structure than those stabilized by paclitaxel, resulting in different biochemical properties (Bonato *et al.* 2025, *in press*).

It also needs to be remembered that the fluorescent probes presented in this article are all derivatives of parent molecules with microtubule-stabilizing or -destabilizing activity, and thus, they exert microtubule-stabilizing or -destabilizing effects themselves. These effects need to be taken into account in order to ensure reliable results in cellular experiments. The optimal use of these molecular probes requires rigorous determination of non-cytotoxic concentration ranges, evaluation of potential off-target effects, and assessment of probe-induced cellular perturbations that may appear in experimental outcomes. The development of fluorescent microtubule-binding probes that exhibit high-affinity target engagement without perturbing polymerization dynamics remains an unmet need, but research on this question is currently ongoing. As we have shown here, there is already a compound (pelophen B) that minimally alters the microtubule lattice; however, this molecule still retains the capacity to modulate microtubule dynamics.²⁵⁶

As a final point, the evolving landscape of microtubule pharmacology asks for continuous advancements in molecular tool development. For example, we are currently witnessing a



Table 1 Table of MTA-derived fluorescent probes^{61,63,70,71,76–81,85–87,89,100,102,104,105,109,110,112,131,133,138,139,141,159,169,170,174,175,178,186,188,194,198,199,210–212,224,225,232,233,240–242,274}

| Name | Structure | Fluorophore | Chemical scaffold and modification site | Binding affinity (M ⁻¹) ^c | Ref. ^d |
|------------------------|------------------------------|------------------------|---|--|-------------------|
| COLCHICINE SITE | | | | | |
| 1a | Colchicine | -- | Colchicine | 1.2 × 10 ⁷ 274 | |
| 1b | MTC | -- | Colchicine ring B | 4.9 × 10 ⁵ 63 | 61 |
| 1c | Lumi-colchicine | Fluorescein | Colchicine C7 | 2.0 × 10 ⁵ 70 4.5 × 10 ⁴ 71 | 70 |
| 1d | Colchicine -504 ^a | BODIPY-504 | Colchicine C7 | 1.7 × 10 ⁵ | 76 |
| 1e | Colchicine -646 ^a | BODIPY-646 | Colchicine C7 | 1.3 × 10 ⁵ | 76 |
| 1f | Dansyl-colchicine | Dansyl | Colchicine C7 | 2.2 × 10 ⁵ | 77 |
| 1g | N-Dansyl-isocolchicine | Dansyl | Isocolchicine C7 | 7.0 × 10 ⁴ | 77 |
| 1h | NBD-colcemid ^a | NBD | Colcemid C7 | 2.0–5.7 × 10 ⁵ | 78 |
| 1i | Tu-SP | Spiropyrane derivative | Colchicine C7 | Not reported | 79 |



Table 1 (continued)

| Name | Structure | Fluorophore | Chemical scaffold and modification site | Binding affinity (M ⁻¹) ^c | Ref. ^d |
|---------------------|---|--------------|---|--|-------------------|
| 1j | Col-BODIPY CB4 | BODIPY | Colchicine C7 | 6.0 × 10 ⁴ | 80 |
| 2a | NSC 613862 | --- | Pteridine | 4.1 × 10 ⁶ | 81 |
| 2b | NSC 613863 | --- | Pteridine | 3.2 × 10 ⁶ | 81 |
| 3a | Morpholino-quinazoline | --- | Quinazoline C2 | Not reported | 85 |
| 4a | CA4 | --- | E-Combretastatin | Not reported | 86 |
| 4b | CA4F | --- | E-Combretastatin | Not reported | 86 |
| 5a | OC9 | --- | Chalcone | 5.9 × 10 ⁷ 88 | 87 |
| 6a | Coumarin-30 | Coumarin-30 | Coumarin | 3.1 × 10 ⁵ | 89 |
| VINCA DOMAIN | | | | | |
| 7a | Vinblastin e-4'-anthranilate ^a | Anthranilate | Vinblastine C20' | 2.5 × 10 ⁴ | 100 |
| 7b | F-VLB | Coumarin | Vinblastine C17 | 7.7 × 10 ⁴ | 102 |
| 7c | BODIPY FL vinblastine ^a | BODIPY FL | Vinblastine C17 | 1.5 × 10 ⁶ 104 | TF |



Table 1 (continued)

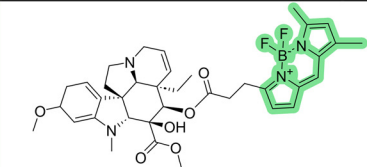
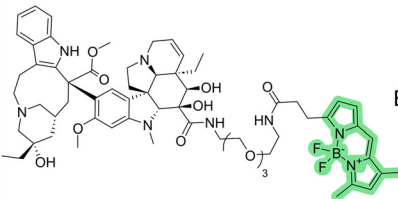
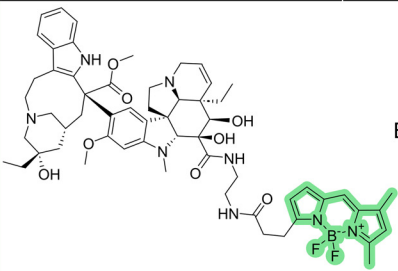
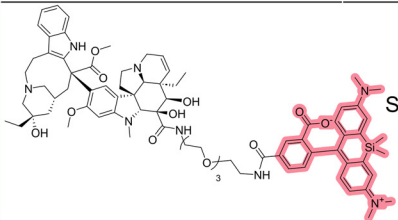
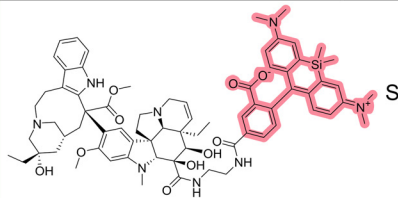
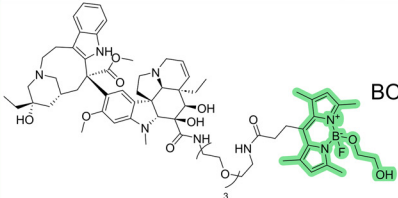
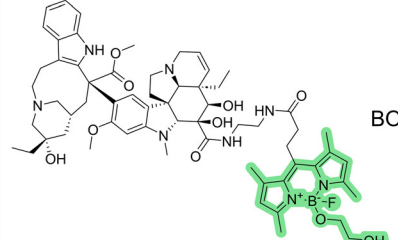
| Name | Structure | Fluorophore | Chemical scaffold and modification site | Binding affinity (M ⁻¹) ^c | Ref. ^d |
|--|---|---------------|---|--|-------------------|
| 7d BODIPY FL vindoline ^a |  | BODIPY FL | Vindoline C17 | 3.9 × 10 ⁶ | 105 |
| 7e BODIPY- PEG ₃ vinblastine |  | BODIPY FL | Vinblastine C16 | Not reported | 109 |
| 7f BODIPY vinblastine |  | BODIPY FL | Vinblastine C16 | Not reported | 109 |
| 7g SiR-PEG ₃ vinblastine |  | SiRhodamine | Vinblastine C16 | Not reported | 109 |
| 7h SiR- vinblastine |  | SiRhodamine | Vinblastine C16 | Not reported | 109 |
| 7i BODIPY- EG-PEG ₃ vinblastine |  | BODIPY(EG) FL | Vinblastine C16 | Not reported | 109 |
| 7j BODIPY-EG vinblastine |  | BODIPY(EG) FL | Vinblastine C16 | Not reported | 109 |



Table 1 (continued)

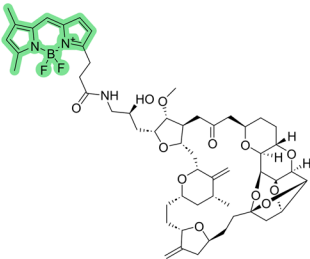
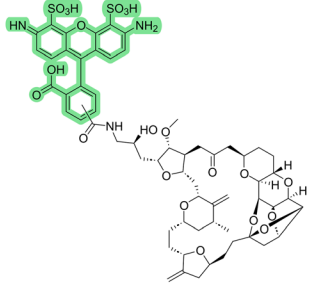
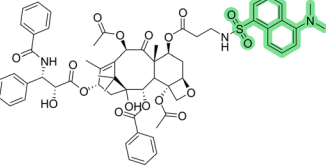
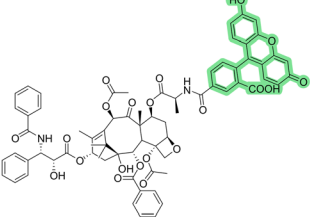
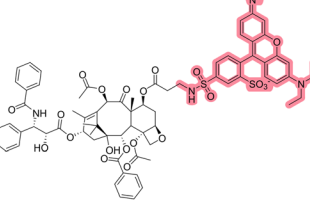
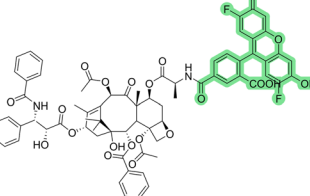
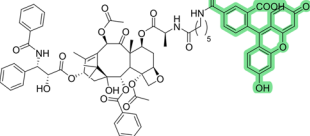
| Name | Structure | Fluorophore | Chemical scaffold and modification site | Binding affinity (M ⁻¹) ^c | Ref. ^d |
|------------------------------------|---|----------------|---|--|-------------------|
| 8a BODIPY-Eribulin |  | BODIPY FL | Eribulin C35 | Not reported | 110 |
| 8b Eribulin488 ^a |  | AlexaFluor 488 | Eribulin C35 | 2.9 × 10 ⁷ | 112 |
| TAXANE SITE | | | | | |
| 9a Dan-PTX |  | Dansyl | Paclitaxel C7 | Not reported | 131 |
| 9b Flutax-1 |  | Fluorescein | Paclitaxel C7 | 2.9 × 10 ⁷ 139 | 133 |
| 9c Rotax |  | Rhodamine | Paclitaxel C7 | Not reported | 138 |
| 9d Flutax-2 ^b |  | Oregon green | Paclitaxel C7 | 2.2 × 10 ⁷ | 139 |
| 9e Hexaflutax |  | Fluorescein | Paclitaxel C7 | 7.2 × 10 ⁵ | 141 |



Table 1 (continued)

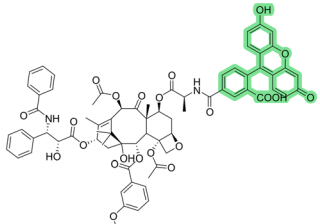
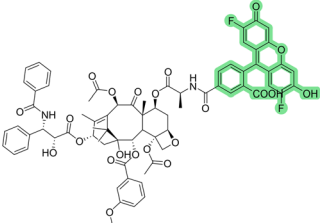
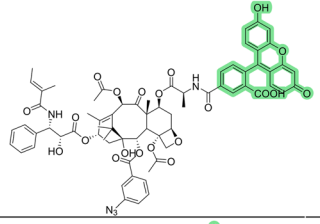
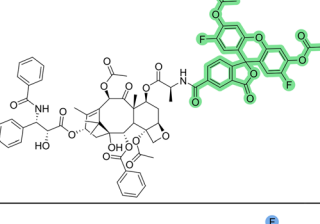
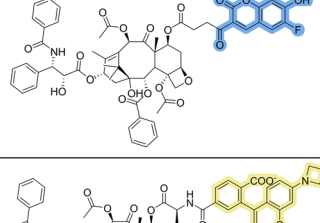
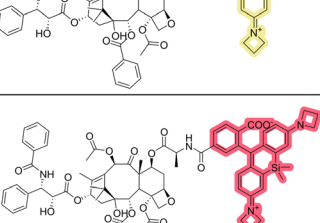
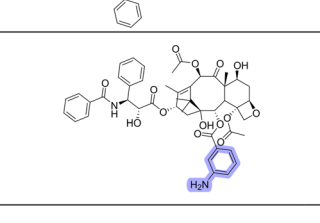

| Name | Structure | Fluorophore | Chemical scaffold and modification site | Binding affinity (M ⁻¹) ^c | Ref. ^d | |
|------|------------------------------------|---|---|--|----------------------------|-----|
| 9f | FChitax-2 |  | Fluorescein | Chitax-11 C7 | 4.4 × 10 ⁸ | 169 |
| 9g | FChitax-3 |  | Oregon green | Chitax-11 C7 | 6.8 × 10 ⁸ | 169 |
| 9h | FChitax-4 |  | Fluorescein | Cephalomane C7 | 1.2 × 10 ⁹ | 169 |
| 9i | Tubulin Tracker Green ^b |  | Oregon green | Paclitaxel C7 | Not reported | TF |
| 9j | PB-Gly-TXL ^a |  | Pacific Blue | Paclitaxel C7 | 2.9 × 10 ⁷ | 170 |
| 9k | JFtaxol549 ^b |  | Janelia Fluor 549 | Paclitaxel C7 | Not reported | T |
| 9l | JFtaxol646 ^b |  | Janelia Fluor 646 | Paclitaxel C7 | 1.1 × 10 ⁶ (SM) | T |
| 9m | 2-AB-PT |  | Anilin | Paclitaxel C2 | 2.0 × 10 ⁷ | 174 |



Table 1 (continued)

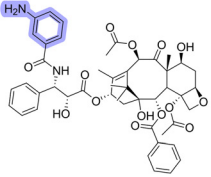
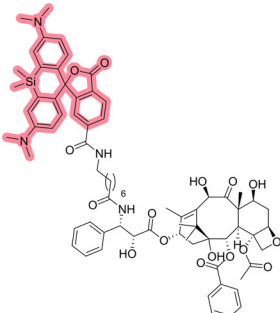
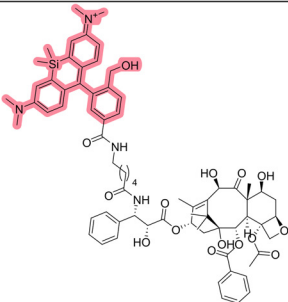
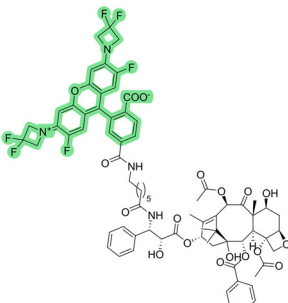
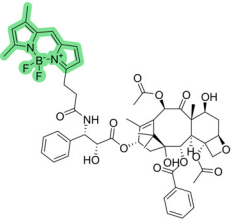
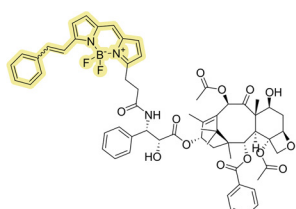
| | Name | Structure | Fluorophore | Chemical scaffold and modification site | Binding affinity (M ⁻¹) ^c | Ref. ^d |
|----|----------------------------|---|-------------------|---|--|-------------------|
| 9n | N-AB-PT ^a |  | Anilin | Paclitaxel C3' | 4.0 × 10 ⁵ | 175 |
| 9o | SiR-tubulin ^b |  | SiRhodamine | Docetaxel C3' | 1.4 × 10 ⁷ (SM) | 178 |
| 9p | HMSiR-tubulin ^a |  | HMSi Rhodamine | Docetaxel C3' | 8.3 × 10 ⁶ | 186 |
| 9q | JFtaxol526 ^b |  | Janelia Fluor 526 | Paclitaxel C3' | Not reported | 188 |
| 9r | BODIPY-FL-PTX |  | BODIPY- FL | Paclitaxel C3' | Not reported | TF |
| 9s | BODIPY 564/570-PTX |  | BODIPY 564/570 | Paclitaxel C3' | Not reported | TF |



Table 1 (continued)

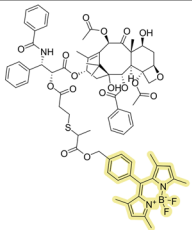
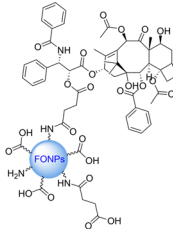
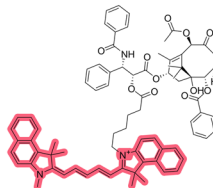
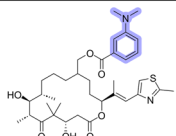
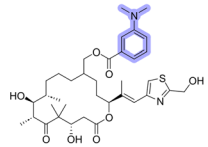
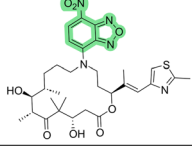
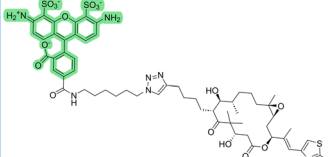
| Name | Structure | Fluorophore | Chemical scaffold and modification site | Binding affinity (M ⁻¹) ^c | Ref. ^d |
|---|---|-------------------------------------|---|--|-------------------|
| 9t PTX-S-BDP |  | BODIPY OH | Paclitaxel C2' | Not reported | 194 |
| 9u PTX-FONPs |  | FONPs | Paclitaxel C2' | Not reported | 198 |
| 9v PTX-Cy5.5 |  | Cy5.5 | Paclitaxel C2' | Not reported | 199 |
| 9w Tubulin Tracker Deep Red ^b | N/A | N/A | N/A | 1.3 × 10 ⁶ (SM) | TF |
| 9x Via Fluor 405/488/647 ^b | N/A | Alexa Fluor, Texas Red, and DyLight | N/A | Not reported | B |
| 9y BioTracker 488 ^b | N/A | N/A | N/A | Not reported | M |
| 9z SPY 555/650/700b -tubulin ^b | N/A | SPY 555/650/700 | N/A | Not reported | S |
| 10a Dimethylam inobenzoyl-epothilone D |  | Dimethylaminoben zoyl | Epothilone D C26 | Not reported | 210 |
| 10b Dimethylam inobenzoyl-C21-hydroxy epothilone D |  | Dimethylaminoben zoyl | Epothilone D C26 | Not reported | 210 |
| 10c NBD-azathilone ^a |  | NBD | Aza-epothilone C12 | 2.2 × 10 ⁶ | 211 |
| 10d GS244 |  | AlexaFluor 488 | Epothilone B C6' | 1.5 × 10 ⁸ 212 | 159 |



Table 1 (continued)

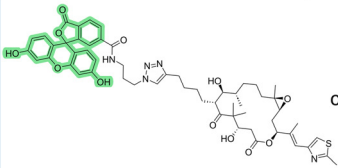
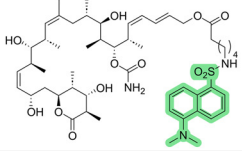
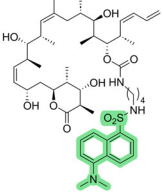
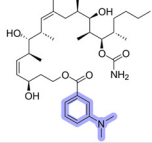
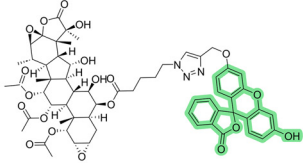
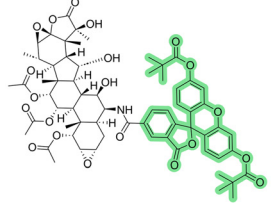
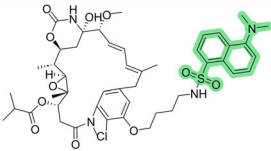
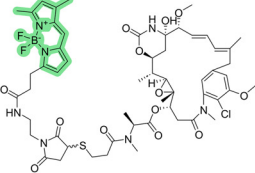
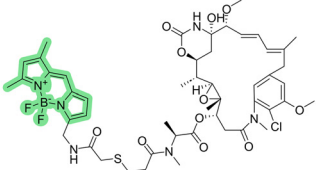
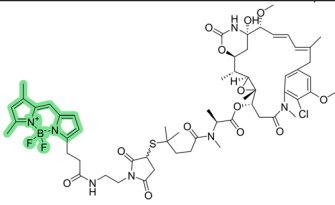
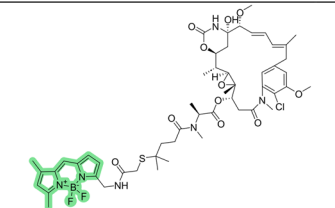
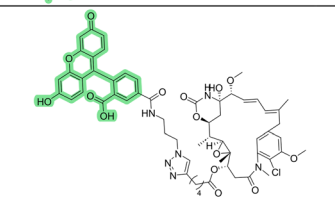
| Name | Structure | Fluorophore | Chemical scaffold and modification site | Binding affinity (M ⁻¹) ^c | Ref. ^d |
|------------------------|---|-------------------------------|---|--|-------------------|
| 10e |  | 6-carboxyfluorescein | Epothilone B C6' | 2.7 × 10 ⁵ | 212 |
| 11a |  | Dansyl | Discodermolide C24 | Not reported | 224 |
| 11b |  | Dansyl | Discodermolide C19 | Not reported | 224 |
| 11c |  | Dimethylaminobenzoic acid | Discodermolide C5 | 4.1 × 10 ⁴ | 225 |
| 12a |  | Fluorescein | Taccalonolide AJ C6 | Not reported | 232 |
| 12b |  | Dipivaloyl-carboxyfluorescein | Taccalonolide AJ C6 | Not reported | 233 |
| MAYTANSINE SITE | | | | | |
| 13a |  | Dansyl | PDM-3 C20 | Not reported | 240 |
| 13b |  | BODIPY X | DM1 C3 | Not reported | 241 |
| 13c |  | BODIPY X | DM1 C3 | Not reported | 241 |



Table 1 (continued)

| Name | Structure | Fluorophore | Chemical scaffold and modification site | Binding affinity (M ⁻¹) ^c | Ref. ^d |
|--------------------------------------|---|-------------|---|--|-------------------|
| 13d DM4-mal-BODIPY |  | BODIPY X | DM4 C3 | Not reported | 241 |
| 13e DM4-BODIPY |  | BODIPY X | DM4 C3 | Not reported | 241 |
| 13f FcMaytansine ^a |  | Fluorescein | Maytansinol C3 | 1.5 × 10 ⁸ | 242 |

^a The binding constant was calculated from the dissociation constant found in the literature following the equation: $K_b = \frac{1}{K_d}$. ^b Commercially available. TF (Thermo-Fisher), T (Tocris), B (Biotium), M (Merck) and S (Spirochrome). ^c References correspond to the publication reporting the binding or dissociation constant when this value is not provided in the probe discovery manuscript. ^d References correspond to the original publication in which the probe was first reported. RGB color codes assigned to represent each fluorescent emission range are as follows: UV (172,172,255); blue (119,173,237); green (126,226,149); yellow (243,237,156); red (252,139,159); and far-red (251,98,124).

paradigm shift in tubulin-directed drug discovery, with a new focus on the tubulin code, tubulin isotype composition and post-translational modifications. This emerging research focus demands not only the identification of isotype-selective ligands²⁷¹ but also their derivatization into fluorescent probes capable of elucidating structure-function relationships within distinct tubulin populations and facilitating the discovery of next-generation therapeutic candidates. Moreover, the generation of probes capable of selectively recognizing microtubules bearing distinct post-translational modifications could revolutionize the field not only from a drug discovery perspective, but also by enabling detailed research into how these modifications regulate microtubule functions, thereby providing a deeper understanding of all aspects of tubulin biology.

The probes discussed in this review provide the foundational framework for the rational design of next-generation tools with potential clinical applications. Future progress will be based on the interplay of continuous advances in imaging technologies, the discovery and tailored synthesis of fluorophores with superior photophysical properties, and the development of safer and more effective drugs, enabled in part by the molecular probes described herein in this review.

Conclusions and expert opinion

Microtubules are highly dynamic cytoskeletal polymers essential for processes such as mitosis, intracellular transport and neuronal homeostasis. Small fluorescent molecules targeting defined binding sites on tubulin and microtubules have become powerful tools to visualize and interrogate microtubule organization and dynamics in living systems, providing unprecedented spatial and temporal resolution and significantly advancing both fundamental and applied research.

Despite their utility, these probes present an intrinsic and often underappreciated limitation: their unavoidable impact on the equilibrium between assembled microtubules and unassembled tubulin implies that small molecules interacting with tubulin cannot be considered passive observers. Because they bind to functionally relevant sites, small fluorescent ligands inherently alter polymerization dynamics and, in the case of stabilizing agents, they can also modify the microtubule lattice structure,²⁷² thereby affecting microtubule-associated protein recognition. Depending on their binding mode, such compounds may promote assembly, suppress catastrophe, or induce depolymerization, meaning that the fluorescent signal itself can perturb the system under observation. From an expert perspective, this dual role, as both the reporter and modulator,



constitutes a fundamental caveat that must be explicitly considered when interpreting experimental data.

Circumventing this limitation is inherently difficult and, in principle, incompatible with microtubule-destabilizing agents. By definition, these compounds act by blocking essential structural transitions, such as the curved-to-straight conformational change of tubulin at the colchicine or gatorbulin sites, or by masking critical interaction surfaces, as do ligands targeting the pironetin, maytansine, or vinca sites. Their use as fluorescent probes therefore unavoidably perturbs the assembly equilibrium. In contrast, microtubule-stabilizing agents provide a more tractable framework for rational probe design. For ligands binding the laulimalide-peloruside site, which stabilize microtubules by bridging adjacent protofilaments, structural perturbations can be minimized through careful selection of the binding moiety of the fluorescent probe. Moreover, recent work from our group has demonstrated that taxane-site ligands can be engineered to be structurally neutral,^{272,273} or by suppressing certain chemical features biologically inactive, by avoiding enforcement of the straight lattice conformation while retaining microtubule binding.¹²⁸

In conclusion, fluorescent small molecules targeting microtubule sites remain invaluable tools, but their use requires a rigorous and critical experimental framework. Future efforts should prioritize the development of probes that minimize perturbation of polymerization equilibrium, together with standardized biophysical and cellular assays to quantify their effects. Only through such careful design and validation can fluorescent microtubule ligands reliably illuminate cytoskeletal biology without introducing unintended functional interference.

Materials and methods

Determination of the binding constants of fluorescent taxanes

Taxane binding sites in stabilized microtubules were prepared as described in a previous study.¹⁴⁰ Flutax-2 was synthesized as described previously.¹³⁹ The binding constants of JFTaxol646, Tubulin Tracker Deep Red and SiR-tubulin (Table 1) were determined by competition with flutax-2 as described previously.¹⁴⁸

Added in proofs

Glauser *et al.* recently reported the synthesis of a novel far-red fluorescent epothilone probe in conjunction with the synthesis of GS244 (**10d**) and their precursor.²⁷⁵ The new conjugate, functionalized with a PEG₃ linker and a Janelia Fluor 646 fluorophore, demonstrates binding to microtubules with an association constant (K_b) of 3.3×10^7 M. Moreover, this probe efficiently labels cellular microtubules, thereby expanding the repertoire of far-red fluorescent taxane-site probes available for advanced microscopy applications.

Author contributions

RP-O, DL-A, VP, and JFD contributed to the concept and design of the manuscript. RP-O and DL-A contributed to the writing.

VP, JFD and KHA critically revised the manuscript and contributed to the writing. All authors contributed to the article and approved the submitted version.

Conflicts of interest

The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

Data availability

No primary research results, software or code have been included and no new data were generated or analysed as part of this review, with the exception of newly determined binding constants for three commercial compounds, which are presented in Table 1.

Acknowledgements

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