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# Effect of fluorine substituents in 4-(1-benzyl-1*H*-benzo[*d*]imidazol-2-yl)thiazole for the study of antiparasitic treatment of cysticercosis on a *Taenia crassiceps* model†

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This work details the synthesis of five *N*-benzylated derivatives of thiabendazoles (**L1–L5**), four of which were previously unreported in the literature (**L2–L5**). The compounds were characterised using a comprehensive array of spectroscopic (FT-IR, <sup>1</sup>H, <sup>13</sup>C{1H}, and <sup>19</sup>F{1H} NMR), spectrometric (MS-EI+) and diffractometric (SC-DRX) techniques. To evaluate the effect of increased fluorine substituents in the *N*-benzyl fragment, we conducted a parasitotoxic activity assay, testing the compounds at various concentrations of unhatched *Taenia crassiceps* cysticerci. The inclusion of the *N*-benzyl fragment and the increase in fluorine substituents led to an enhancement in the lipophilicity of thiabendazoles.

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## Introduction

Benzimidazoles are bicyclic heteroaromatic compounds that have attracted the scientific community's attention since their introduction as antiparasitic agents in the 1960s.<sup>1,2</sup> These heterocyclic pharmacophores serve as essential scaffolds in designing new molecules with biological activity or clinical application. Benzimidazoles are among the ten most used nitrogenous heterocycles in medicines approved by the U.S. Food and Drug Administration (FDA), both in human and animal medicine, due to their notable pharmacological

properties.<sup>3,4</sup> These compounds are capable of accepting or donating protons, allowing the formation of weak interactions, which promotes the union of the therapeutic targets with these heterocycles. Structure–activity relationship (SAR) analyses indicate that the placement of substituents in the N1, C2, C5 or C6 positions influences their therapeutic activity.<sup>5,6</sup> Modifications in benzimidazoles affect their binding capacity with proteins since new weak interactions and new electronic and steric effects arise, which alter their ability to recognise specific enzymes and modify their pharmacological potential.<sup>7,8</sup> These molecules have a structure similar to those of the nitrogenous bases known as purines (adenine and guanine), which facilitates their application in various biological activities, such as analgesic, anticancer, anticoagulant, anticonvulsant, antidiabetic, antihypertensive, antihistaminic, anti-inflammatory, antimalarial, antimicrobial, antioxidant, antipsychotic, antituberculosis, antitumor, antiulcer, antiviral, and contraceptive activities, among others.<sup>9–11</sup>

One of the best-known benzimidazoles is 4-(1*H*-benzo[*d*]imidazol-2-yl)thiazole, commonly called thiabendazole (TBZ). This compound inhibits the helminth-specific mitochondrial enzyme (fumarate reductase), which is why it is used as an anthelmintic agent in animals and humans.<sup>12,13</sup> TBZ was the first benzimidazole-based drug approved by the FDA for treating human gastrointestinal parasites. Even so, it was later withdrawn from the market because other anthelmintic agents with better tolerance emerged.<sup>14,15</sup> Nonetheless, this drug is still available in many countries and is mainly used in veteri-

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ary medicine. TBZ is also used as a fungicide and an anthelmintic in fruits, vegetables, and various food products.<sup>16,17</sup> This compound generally has low toxicity; however, the U.S. Environmental Protection Agency (EPA) has classified it as a probable carcinogen at doses high enough to cause disturbances in thyroid hormone balance.<sup>18,19</sup>

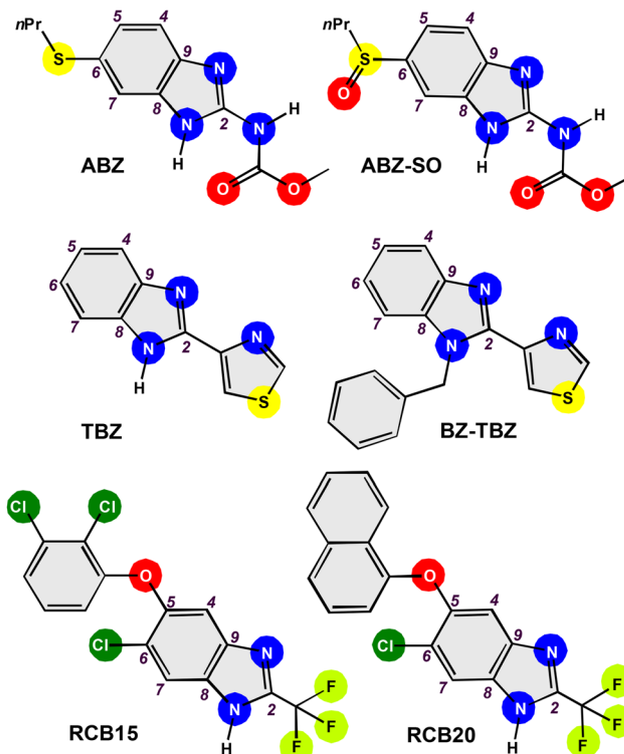
Brown and co-workers (1961) reported the preparation of 4-(1-benzyl-1*H*-benzo[*d*]imidazol-2-yl)thiazole (BZ-TBZ) when searching for a new class of broad-spectrum anthelmintic agents against gastrointestinal parasites of domestic animals. For this, TBZ was functionalised in the N1 position using benzoyl iodide.<sup>20</sup> Several studies have been reported on synthesising and characterising this TBZ derivative.<sup>21–23</sup> Even SAR experimental studies have been reported on its antimicrobial, anti-inflammatory and antiangiogenic activities, and as a vascular disruptor and methionine aminopeptidase inhibitor.<sup>24–29</sup>

On the other hand, *Taenia crassiceps* is a cestode found in wild canids and rodents. This parasite presents metabolic and antigenic similarities to different species of the *Taeniidae* family, such as *Taenia solium*, which makes it useful as an experimental model in cysticercosis studies.<sup>30,31</sup> Human cysticercosis caused by *Taenia crassiceps* is unusual; however, isolated cases have been reported in immunocompromised and immunocompetent patients.<sup>32,33</sup> One of the primary drugs used in the treatment of this disease is albendazole (ABZ), a derivative of benzimidazole. Nonetheless, parasite resistance to this drug has been reported, which has motivated the search for new alternatives for treating cysticercosis.<sup>34,35</sup> Thus, developing new benzimidazole derivatives aims to increase antiparasitic activity by slightly modifying traditional anthelmintics.<sup>36,37</sup> New compounds have been synthesised from 2-(trifluoromethyl)-1*H*-benzo[*d*]imidazole, presenting improved antiparasitic activity. For instance, 6-chloro-5-(2,3-dichlorophenoxy)-2-(trifluoromethyl)-1*H*-benzo[*d*]imidazole (RCB15) and 6-chloro-5-(naphthalen-1-yloxy)-2-(trifluoromethyl)-1*H*-benzo[*d*]imidazole (RCB20) have been chosen as promising molecules for the treatment of this disease, Scheme 1. These fluorinated benzimidazole derivatives had a greater effect on *Taenia crassiceps* cysticerci than ABZ and albendazole sulfoxide (ABZ-SO).<sup>38,39</sup> Therefore, this work reports the synthesis and characterisation of BZ-TBZ derivatives with different degrees of fluorination in the *N*-benzylic arm to study their effect on antiparasitic activity.

## Materials and methods

### Experimental details

All starting materials were purchased from commercial suppliers and used without further purification. The reagents employed were thiabendazole, sodium hydride 90%, benzyl bromide 98%, 3-fluorobenzyl chloride 96%, potassium iodide 99%, 2,3,6-trifluorobenzyl bromide 97%, 2,3,4,5,6-pentafluorobenzyl bromide 99%, and 4-(trifluoromethyl)benzyl bromide 98%. Melting points were determined directly using a



Scheme 1 Benzimidazole derivatives with anticysticercosis activity.

Fisherbrand™ digital apparatus (open capillary). Infrared spectra were obtained using an FT-IR Bruker Alpha ATR spectrophotometer. The <sup>1</sup>H, <sup>13</sup>C{<sup>1</sup>H} and <sup>19</sup>F{<sup>1</sup>H} NMR experiments were carried out by using Bruker Avance III 300 MHz equipment using CDCl<sub>3</sub> in 5 mm tubes, and subsequently, the spectra were processed using MestReNova® software. Chemical shift values (δ) are reported in parts per million (ppm) relative to the internal reference (TMS). Mass spectra were recorded using a Jeol JSM-SX102A spectrometer at 10 kV in the EI+ mode. The colourless prismatic crystals were placed in a Bruker Smart Apex II X-ray diffractometer with a molybdenum X-ray source (λ = 0.71073 Å). The structures were solved and refined using Patterson's methods with the programs SHELXS-2014/7 and SHELXL-2019/2.<sup>40</sup> Molecular structures and supramolecular arrangements were analysed using Mercury® software. Additional information on structural data collection, resolution, and refinement can be found in the ESI.† The three structural files were deposited at the Cambridge Crystallographic Data Center (CCDC). The assigned deposit numbers for L1, L2, and L4 were 2350921, 2350919, and 2350920,† respectively. Table 1 contains the crystallographic parameters, and Tables S1–S3† contain the distances and bond angles.

### Synthesis of *N*-benzylated compounds

The compounds (L1–L5) were synthesised from the deprotonation of thiabendazoles (1 mmol, 0.2013 g) in the presence of sodium hydride (1 mmol, 0.0222 g). Subsequently, the product



Table 1 Crystal data and structure refinement for L1, L2 and L4

|   | L1   | L2   | L4   |
|---|--|--|--|
| Empirical formula                                   | C <sub>17</sub> H <sub>13</sub> N <sub>3</sub> S   | C <sub>17</sub> H <sub>12</sub> FN <sub>3</sub> S  | C <sub>17</sub> H <sub>8</sub> F <sub>5</sub> N <sub>3</sub> S   |
| Formula weight                                      | 291.36   | 309.36   | 381.32   |
| Temperature   | 298(2) K   | 298(2) K   | 298(2) K   |
| Wavelength  | 0.71073 Å  | 0.71073 Å  | 0.71073 Å  |
| Monoclinic crystal system                           | Monoclinic   | Monoclinic   | Monoclinic   |
| Space group   | P2 <sub>1</sub> /n   | P2 <sub>1</sub> /n   | P2 <sub>1</sub> /c   |
| Unit cell dimensions                                | <i>a</i> = 17.3372(8), <i>α</i> = 90°<br><i>b</i> = 9.2800(4), <i>β</i> = 113.263(2)°<br><i>c</i> = 19.3163(8), <i>γ</i> = 90° | <i>a</i> = 13.1322(4) Å, <i>α</i> = 90°<br><i>b</i> = 6.0454(2) Å, <i>β</i> = 107.2870(10)°<br><i>c</i> = 19.1953(7) Å, <i>γ</i> = 90° | <i>a</i> = 17.173(4) Å, <i>α</i> = 90°<br><i>b</i> = 8.1616(18) Å, <i>β</i> = 93.871(5)°<br><i>c</i> = 10.976(3) Å, <i>γ</i> = 90° |
| Volume  | 2855.1(2) Å <sup>3</sup>   | 1455.07(8) Å <sup>3</sup>  | 1534.8(6) Å <sup>3</sup>   |
| Z   | 8  | 4  | 4  |
| Density (calculated)                                | 1.356 Mg m <sup>-3</sup>   | 1.412 Mg m <sup>-3</sup>   | 1.650 Mg m <sup>-3</sup>   |
| Absorption coefficient                              | 0.223 mm <sup>-1</sup>   | 0.232 mm <sup>-1</sup>   | 0.273 mm <sup>-1</sup>   |
| <i>F</i> (000)                                      | 1216   | 640  | 768  |
| Crystal size  | 0.387 × 0.243 × 0.066 mm <sup>3</sup>  | 0.462 × 0.167 × 0.090 mm <sup>3</sup>  | 0.280 × 0.180 × 0.150 mm <sup>3</sup>  |
| Theta range for data collection                     | 2.028 to 25.328°   | 1.673 to 25.390°   | 2.377 to 25.472°   |
| Index ranges  | −20 ≤ <i>h</i> ≤ 20, −11 ≤ <i>k</i> ≤ 10,<br>−23 ≤ <i>l</i> ≤ 22   | −15 ≤ <i>h</i> ≤ 15, −7 ≤ <i>k</i> ≤ 7,<br>−23 ≤ <i>l</i> ≤ 23   | −20 ≤ <i>h</i> ≤ 20, −9 ≤ <i>k</i> ≤ 9,<br>−13 ≤ <i>l</i> ≤ 13   |
| Reflections collected                               | 25 355   | 12 341   | 13 161   |
| Independent reflections                             | 5207 [ <i>R</i> (int) = 0.0919]  | 2674 [ <i>R</i> (int) = 0.0421]  | 2837 [ <i>R</i> (int) = 0.0851]  |
| Completeness to theta                               | 25.242°, 99.9%   | 25.242°, 100.0%  | 25.242°, 100.0%  |
| Absorption correction                               | None   | None   | None   |
| Refinement method                                   | Full-matrix least-squares on <i>F</i> <sup>2</sup>   | Full-matrix least-squares on <i>F</i> <sup>2</sup>   | Full-matrix least-squares on <i>F</i> <sup>2</sup>   |
| Data/restraints/parameters                          | 5207/0/379   | 2674/0/199   | 2837/608/357   |
| Goodness-of-fit on <i>F</i> <sup>2</sup>            | 0.903  | 0.967  | 0.958  |
| Final <i>R</i> indices [ <i>I</i> > 2σ( <i>I</i> )] | <i>R</i> <sub>1</sub> = 0.0566, <i>wR</i> <sub>2</sub> = 0.1353  | <i>R</i> <sub>1</sub> = 0.0344, <i>wR</i> <sub>2</sub> = 0.0791  | <i>R</i> <sub>1</sub> = 0.0501, <i>wR</i> <sub>2</sub> = 0.1121  |
| <i>R</i> indices (all data)                         | <i>R</i> <sub>1</sub> = 0.1475, <i>wR</i> <sub>2</sub> = 0.1725  | <i>R</i> <sub>1</sub> = 0.0545, <i>wR</i> <sub>2</sub> = 0.0885  | <i>R</i> <sub>1</sub> = 0.1201, <i>wR</i> <sub>2</sub> = 0.1392  |
| Largest diff. peak and hole                         | 0.354 and −0.397 e Å <sup>-3</sup>   | 0.122 and −0.277 e Å <sup>-3</sup>   | 0.164 and −0.246 e Å <sup>-3</sup>   |

was reacted with each fluoro-substituted benzyl halide under a nitrogen atmosphere at the reflux of anhydrous tetrahydrofurans for 24 h. L1 and L3–L5 were obtained from fluoro-substituted benzyl bromides, while L2 was obtained from 3-fluorobenzyl chloride and potassium iodide. The reactions were monitored by thin-layer chromatography (TLC) in a dichloromethane–methanol (95 : 5) mixture and developed using a UV lamp. Afterwards, the reaction products were purified by column chromatography and concentrated to dryness using a rotary evaporator connected to a high vacuum. L1, L2 and L4 crystallised from the slow evaporation of a mixture of dichloromethane with acetone.

**4-(1-Benzyl-1*H*-benzo[*d*]imidazol-2-yl)thiazole (L1).** It was synthesised from benzyl bromide (1 mmol, 0.1745 g). Light brown solid (0.2244 g, 77%). M.p. 146–148 °C. IR (ATR, *ν* cm<sup>-1</sup>): 3111 (w), 3076 (w), 2916 (w), 1602 (w), 1450 (m), 1397 (m), 1303 (m), 1283 (w), 901 (m), 880 (m), 830 (m), 734 (s). <sup>1</sup>H NMR (300 MHz, CDCl<sub>3</sub>, *δ*, ppm): 8.75 (s, 1H), 8.29 (s, 1H), 7.75 (d, 1H), 7.23–7.04 (m, 6H), 5.98 (s, 2H). <sup>13</sup>C{<sup>1</sup>H} NMR (75 MHz, CDCl<sub>3</sub>, *δ*, ppm): 153.07, 147.83, 146.96, 143.09, 137.11, 136.01, 128.71, 127.53, 126.74, 123.31, 122.87, 121.38, 119.79, 110.70, 48.54. MS-EI+ (*m/z*): 291, 214, 91.

**4-(1-(3-Fluorobenzyl)-1*H*-benzo[*d*]imidazol-2-yl)thiazole (L2).** It was synthesised from 3-fluorobenzyl chloride (1 mmol, 0.1506 g) and potassium iodide (1.5 mmol, 0.2515 g). White solid (0.2413 g, 78%). M.p. 140–142 °C. IR (ATR, *ν* cm<sup>-1</sup>): 3122 (w), 3080 (w), 1614 (m), 1586 (m), 1480 (m), 1449 (m), 1437 (m), 1352 (w), 1295 (w), 1253 (w), 1141 (w), 939 (m), 829 (w), 746 (s). <sup>1</sup>H NMR (300 MHz, CDCl<sub>3</sub>, *δ*, ppm): 8.78 (s, 1H), 8.27

(s, 1H), 7.75 (d, 1H), 7.25–7.09 (m, 4H), 6.86–6.76 (m, 3H), 5.98 (s, 2H). <sup>19</sup>F{<sup>1</sup>H} NMR (282.77 MHz, CDCl<sub>3</sub>, *δ*, ppm): −112.46. MS-EI+ (*m/z*): 309, 308, 214, 195, 135, 109.

**4-(1-(2,3,6-Trifluorobenzyl)-1*H*-benzo[*d*]imidazol-2-yl)thiazole (L3).** It was synthesised from trifluorobenzyl bromide (1 mmol, 0.2320 g). White solid (0.2590 g, 75%). M.p. 166–168 °C. IR (ATR, *ν* cm<sup>-1</sup>): 3130 (w), 3030 (w), 3004 (w), 1493 (s), 1306 (m), 1242 (s), 1037 (m), 879 (w), 812 (m), 733 (s). <sup>1</sup>H NMR (300 MHz, CDCl<sub>3</sub>, *δ*, ppm): 8.87 (s, 1H), 8.60 (s, 1H), 7.78 (d, 1H), 7.34 (d, 1H), 7.24 (dt, 2H), 6.97 (m, 1H), 6.71 (t, 1H), 6.34 (s, 2H). <sup>19</sup>F{<sup>1</sup>H} NMR (282.77 MHz, CDCl<sub>3</sub>, *δ*, ppm): −118.77, −136.53, −141.27. MS-EI+ (*m/z*): 345, 326, 214, 145.

**4-(1-(Perfluorophenyl)methyl)-1*H*-benzo[*d*]imidazol-2-yl)thiazole (L4).** It was synthesised from 2,3,4,5,6-pentafluorobenzyl bromide (1 mmol, 0.2636 g). White solid (0.2669 g, 70%). M.p. 150–152 °C. IR (ATR, *ν* cm<sup>-1</sup>): 3122 (w), 3070 (w), 3002 (w), 2968 (w), 1657 (w), 1502 (s), 1400 (m), 1306 (m), 1120 (m), 948 (m), 821 (m), 732 (s). <sup>1</sup>H NMR (300 MHz, CDCl<sub>3</sub>, *δ*, ppm): 8.85 (s, 1H), 8.50 (s, 1H), 7.77 (d, 1H), 7.28–7.19 (m, 3H), 6.33 (s, 2H). <sup>19</sup>F{<sup>1</sup>H} NMR (282.77 MHz, CDCl<sub>3</sub>, *δ*, ppm): −141.84, −153.09, −161.02. MS-EI+ (*m/z*): 381, 362, 303, 214, 181.

**4-(1-(4-(Trifluoromethyl)benzyl)-1*H*-benzo[*d*]imidazol-2-yl)thiazole (L5).** It was synthesised from 4-(trifluoromethyl)benzyl bromide (1 mmol, 0.2439 g). White solid (0.2803 g, 78%). M.p. 144–146 °C. IR (ATR, *ν* cm<sup>-1</sup>): 3119 (w), 1619 (w), 1401 (s), 1100 (s), 1063 (s), 911 (m), 817 (m), 738 (s). <sup>1</sup>H NMR (300 MHz, CDCl<sub>3</sub>, *δ*, ppm): 8.78 (s, 1H), 8.41 (s, 1H), 7.79 (d, 1H), 7.44 (d, 2H), 7.25 (m, 3H), 7.19 (d, 2H), 6.08 (s, 2H). <sup>13</sup>C{<sup>1</sup>H} NMR (75 MHz, CDCl<sub>3</sub>, *δ*, ppm): 153.16, 147.61, 146.79, 143.07,



141.21, 135.77, 126.94, 125.72, 123.54, 123.12, 121.56, 119.95, 110.28, 48.19.  $^{19}\text{F}\{^1\text{H}\}$  NMR (282.77 MHz,  $\text{CDCl}_3$ ,  $\delta$ , ppm):  $-62.55$ . MS-EI+ ( $m/z$ ): 359, 358, 214, 159.

### Parasite viability *in vitro*

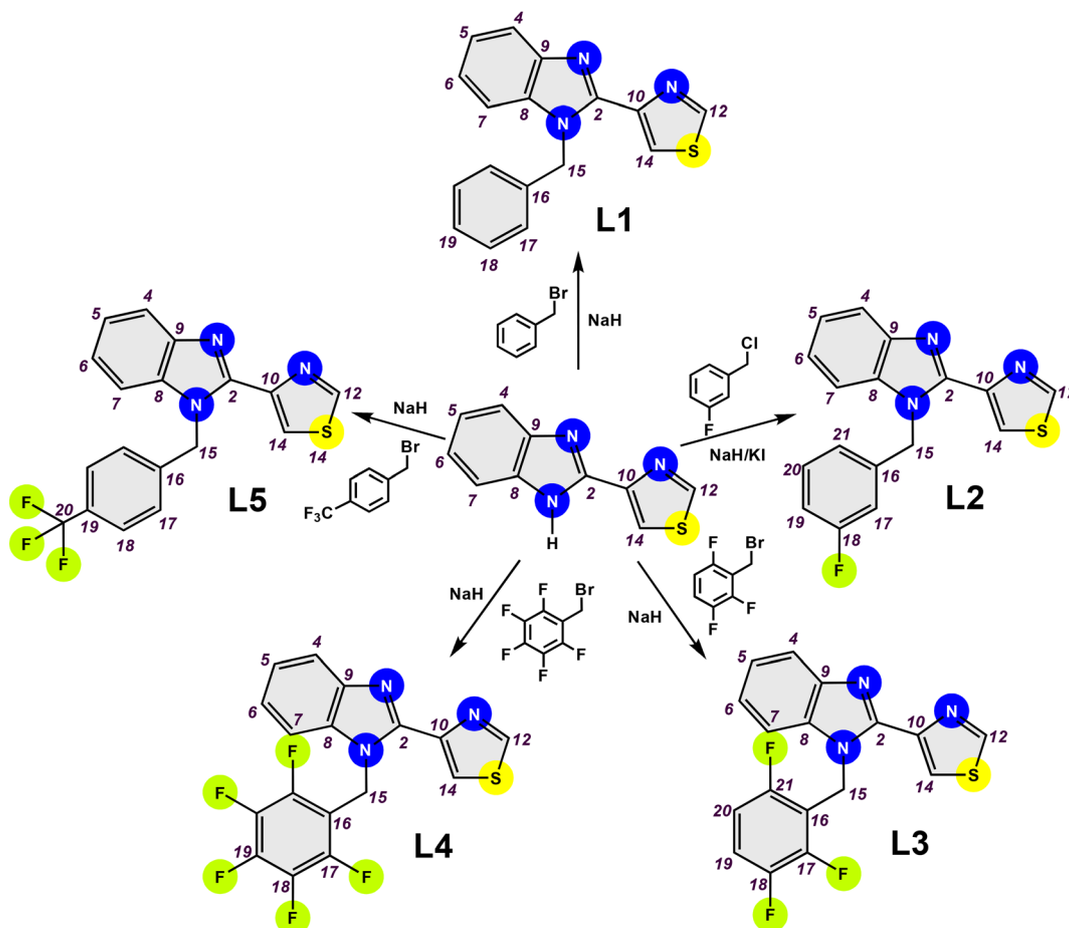
*Taenia crassiceps* strain Wake Forest University (WFU) cysticerci was obtained from the peritoneum of female BALB/cAnN mice at five months of intraperitoneal infection. Cysticerci were washed with sterile phosphate-buffered saline (PBS) and used for *in vitro* assays. The effect of parasitotoxic activity of L1–L5 was determined on the viability and integrity of *Taenia crassiceps* cysticerci. To do this, seven groups of twenty parasites were incubated in microplates using Roswell Park Memorial Institute (RPMI) culture medium at 37 °C in 0.5%  $\text{CO}_2$  for 24 h and analysed at 3, 6 and 24 h. Viability was evaluated with four different concentrations of the compounds (100, 200, 300, and 500  $\mu\text{M}$ ) compared to the same concentrations for the reference drugs, ABZ and TBZ. Subsequently, the parasites were incubated for 1 h in pig bile diluted at a 1 : 3 ratio with RPMI to observe evagination. Viability was estimated by evagination, contractile movements and damage to the bladder wall of the cysticerci, all observed under a Nikon Eclipse TS100 inverted microscope.

## Results and discussion

### Synthesis and structural characterisation

The *N*-benzylation reaction, a crucial step in our research, was performed at the N1 position of TBZ. This stage is fundamental in enhancing the lipophilicity and bioavailability of the new compounds, a key aspect of our study. This reaction also allowed the introduction of fluorine substituents on the *N*-benzylic arm in four of the five compounds. Therefore, L1 and L3–L5 were prepared by one-step synthesis, which consisted of reacting TBZ with different fluorinated benzyl bromides in the presence of sodium hydride under a nitrogen atmosphere. In the case of L2, 3-fluorobenzyl chloride was used in the presence of potassium iodide under the aforementioned conditions, Scheme 2. Most of the compounds recovered as white solids (L2–L5), except for L1, which was light brown. The compounds were obtained with yields between 70 and 78% and showed good solubility in common organic solvents, such as methanol, ethanol, acetone, dichloromethane, chloroform and dimethyl sulfoxide.

IR spectra provided the first evidence of the formation of the *N*-benzylated compounds. The absence of the signal due to the C–Br bond ( $\nu = 650\text{--}515\text{ cm}^{-1}$ ) from benzyl bromides indi-



Scheme 2 General procedure for the preparation of fluorinated compounds derived from TBZ.



cated that the reaction had been completed. Besides, the absence of overtones in the spectra of each ligand ( $\nu = 2800\text{--}2700\text{ cm}^{-1}$ ) is an expected behaviour due to the lack of prototropic tautomerism of benzimidazole. The predictable peaks corresponding to the C=N, C-N and C-S bonds appeared at  $\nu = 1657\text{--}1602\text{ cm}^{-1}$ ,  $1357\text{--}1310\text{ cm}^{-1}$  and  $1141\text{--}1037\text{ cm}^{-1}$ , respectively, while the signal due to the C-F bond of the fluorinated rings in **L2**–**L4** was detected at  $\nu = 948\text{--}911\text{ cm}^{-1}$ . The trifluoromethyl group in **L5** was detected at  $\nu = 1100\text{ cm}^{-1}$ . In addition, the bands due to the aromatic C-H bond were observed at  $\nu = 3130\text{--}3000\text{ cm}^{-1}$ , Fig. S1–S5.†

$^1\text{H}$  NMR spectra showed evidence of *N*-benzylation, with the methylene signal assigned to the range of  $\delta = 6.34\text{--}5.98\text{ ppm}$  (H15). For **L3** and **L4**, this same signal shifted to higher frequencies due to the greater number of fluorine substituents on the *N*-benzylic arm. The signals of thiazole rings appeared in the range of  $\delta = 8.87\text{--}8.78\text{ ppm}$  (H12) and  $\delta = 8.60\text{--}8.27\text{ ppm}$  (H14). The signals of the benzimidazole rings (H4–H7) and the *N*-benzyl fragment (H17–H21) overlapped in the aromatic region of  $\delta = 7.80\text{--}6.68\text{ ppm}$ . In all cases, the H4 signal corresponds to a doublet in the range of  $\delta = 7.80\text{--}7.74\text{ ppm}$  due to the absence of prototropic tautomerism in benzimidazole, Fig. S6–S10.†

In the  $^{13}\text{C}\{^1\text{H}\}$  NMR spectra of **L1** and **L5**, the methylene carbon (C15) signal was assigned at 48.5 ppm for **L1** and 48.2 ppm for **L5**. The most displaced carbon in both compounds was the thiazole carbon (C12) due to its proximity to the nitrogen and sulphur heteroatoms (N11 and S13), at 153.1 ppm for **L1** and 153.2 ppm for **L5**. One of the most significant changes was observed in the quaternary benzyl carbon (C16) because it is found between the nitrogen atoms (N1 and N3), shifting from 137.1 ppm for **L1** to 141.2 ppm for **L5**. The coupling between  $^{19}\text{F}$  and  $^{13}\text{C}$  allowed the signals of the trifluoromethyl group to be assigned at 125.7 ppm (C20) and 129.6 ppm (C19), Fig. S11 and S12.†

Additionally, **L2**–**L5** were characterised by  $^{19}\text{F}\{^1\text{H}\}$  NMR. The observed signals were consistent with the molecular structures proposed for the four fluorinated compounds. **L2** and **L5** presented a single signal. In the case of **L2**, the signal was observed at  $-112.5\text{ ppm}$  and corresponded to the fluorine atom placed in the *meta* position of the benzyl ring. In the case of **L5**, the signal was observed at  $-62.8\text{ ppm}$  and corresponded to the trifluoromethyl group placed in the *para* position of the benzyl ring. **L3** and **L4** exhibited the signals corresponding to the position of the fluorine atoms around the benzyl ring. For example, at  $-118.7$ ,  $-136.5$ , and  $-141.3\text{ ppm}$  for **L3**, which correspond to the fluorine atoms placed at C17, C18 and C21, and at  $-141.8$ ,  $-153.1$  and  $-161.0\text{ ppm}$  for **L4**, which correspond to the fluorine atoms placed at C17, C19 and C18, respectively, Fig. S13–S16.† In the MS-EI+ data of **L1**–**L5**, the peak corresponding to the molecular ion of the five compounds was identified at  $m/z = 291$ ,  $309$ ,  $345$ ,  $381$ ,  $359$ . In addition, the fragment corresponding to TBZ was found at  $m/z = 214$  for all compounds, Fig. S17–S21.†

### Crystallographic analysis

The molecular structures of **L1**, **L2** and **L4** with an ellipsoid representation at a 50% probability level are presented in

Fig. 1. **L1** presents two independent molecules in the asymmetric unit, crystallised in a monoclinic system with a  $P21/n$  space group with eight molecules in the unit cell. It is important to note that the benzimidazole ring approaches coplanarity with the thiazole ring with interplane angles of  $9.68^\circ$  and  $14.91^\circ$  between both molecules. In addition, the *N*-benzyl fragment has angles between planes of  $83.95^\circ$  and  $88.01^\circ$  between both molecules. On the other hand, **L2** crystallised in a monoclinic system with a  $P21/n$  space group with four molecules in the unit cell. Unlike **L1**, this compound crystallised with one molecule in the asymmetric unit. Furthermore, the presence of a fluorine atom in the meta position of the *N*-benzyl fragment can be observed. Finally, **L4** crystallised into a monoclinic system with a  $P21/c$  space with four molecules in the unit cell and one in the asymmetric unit. The N1 nitrogen atom bonds to the pentafluorobenzyl ring with an interplane angle of  $74.20^\circ$  concerning the benzimidazole ring. This compound presents disorder in the benzimidazole and thiazole rings.

In **L1**, the C20–H20...N3 hydrogen bond interactions stand out. These interactions can generate a 1D arrangement with an H20...N3 interaction distance of  $2.60\text{ \AA}$  ( $\sum r_{\text{vdw}} = 2.75\text{ \AA}$ ) and a bond angle  $\angle\text{C20–H20–N3}$  of  $141^\circ$ . The two molecules in the asymmetric unit are related by the interactions C25–H25...Cg1 and C31–H31...Cg1, forming a significant part of the molecular structure. The distances between the hydrogen atoms and the Cg1 centroid of the C16–C21 rings are  $2.84$  and  $2.80\text{ \AA}$ , respectively. The C–H... $\pi$  interactions and the C20–H20...N3 hydrogen bond, perpendicular to each other, form a laminar arrangement parallel to the *ac* plane, Fig. 2. In **L2**, the N3 nitrogen atom of benzimidazole forms two C–H...N interactions, giving rise to a 1D arrangement along the *b*-axis. These arrangements present a distance of  $2.55$  and  $2.74\text{ \AA}$  ( $\sum r_{\text{vdw}} = 2.75\text{ \AA}$ ) for C15–H15B...N3 and C14–H14...N3, respectively. The observed angle for  $\angle\text{C15–H15B–N3}$  is  $174^\circ$ , and for the  $\angle\text{C14–H14–N3}$  is  $149^\circ$ . In this same supramolecular arrangement, a C4–H4...S10 interaction is observed at a distance of  $2.87\text{ \AA}$  ( $\sum r_{\text{vdw}} = 3.00\text{ \AA}$ ) with an angle  $\angle\text{C4–H4–S10}$  of  $156^\circ$ . The arrangement is complemented by a  $\pi$ ... $\pi$  interaction between the benzimidazole's six-membered ring and the thiazole's five-membered ring with a distance of  $3.772\text{ \AA}$  between centroids. The complexity of the molecular arrangements is showcased in the 2D and 3D arrangements, which are stabilised by C–H... $\pi$  interactions. For instance, the C19–H19...Cg1 interaction, together with the C–H...N interactions, generates a 2D arrangement parallel to the *bc* plane, Fig. 3. In **L4**,  $\pi$ ... $\pi$  interactions were found between the pentafluorinated system and the thiazole ring. The distance between the centroids was  $3.74\text{ \AA}$ , and these interactions stabilise the 3D supramolecular arrangement, Fig. 4.

### Antiparasitic activity

Since 1979, when treatment with mebendazole (MBZ) at a single dose ( $50\text{ mg kg}^{-1}$  in rats) was shown to be effective in controlling infections caused by *Taenia crassiceps*,<sup>41</sup> multiple studies have been conducted to evaluate the effect of traditional drugs in the treatment of this disease. For example, it was proposed that combining praziquantel (PZQ) and ABZ-SO



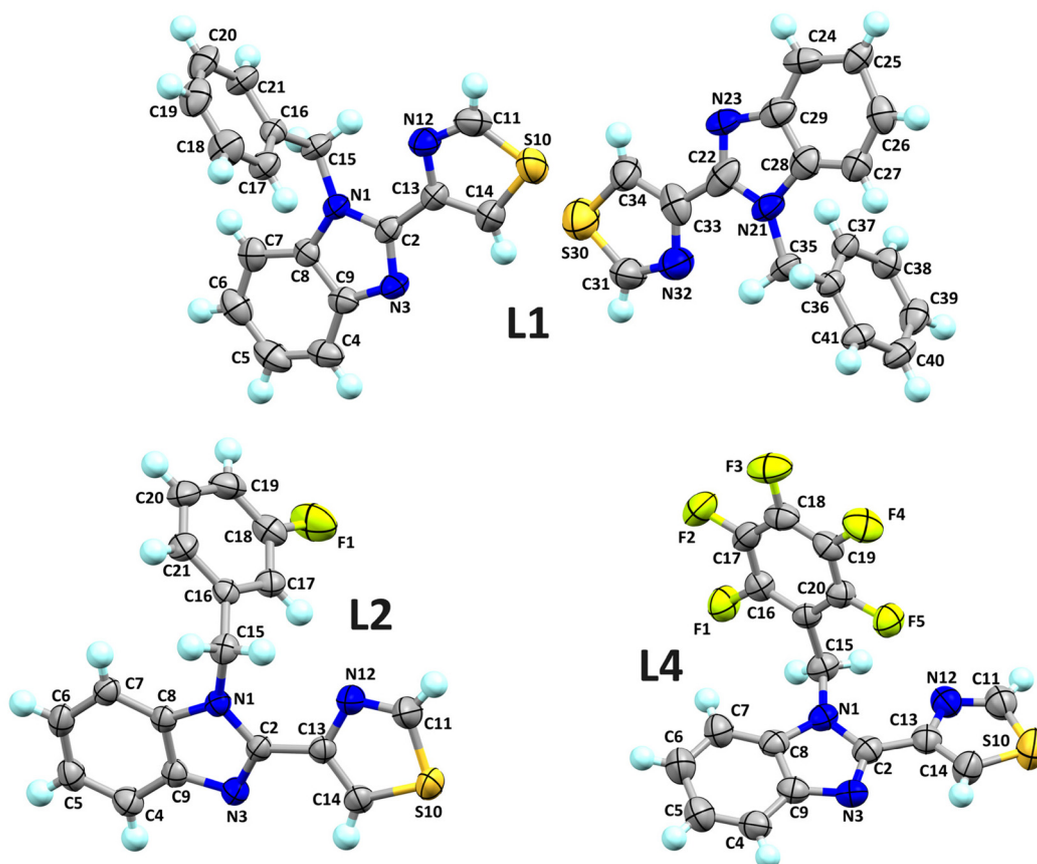


Fig. 1 Ellipsoidal representation at a 50% probability level of the molecular structures of L1, L2 and L4.

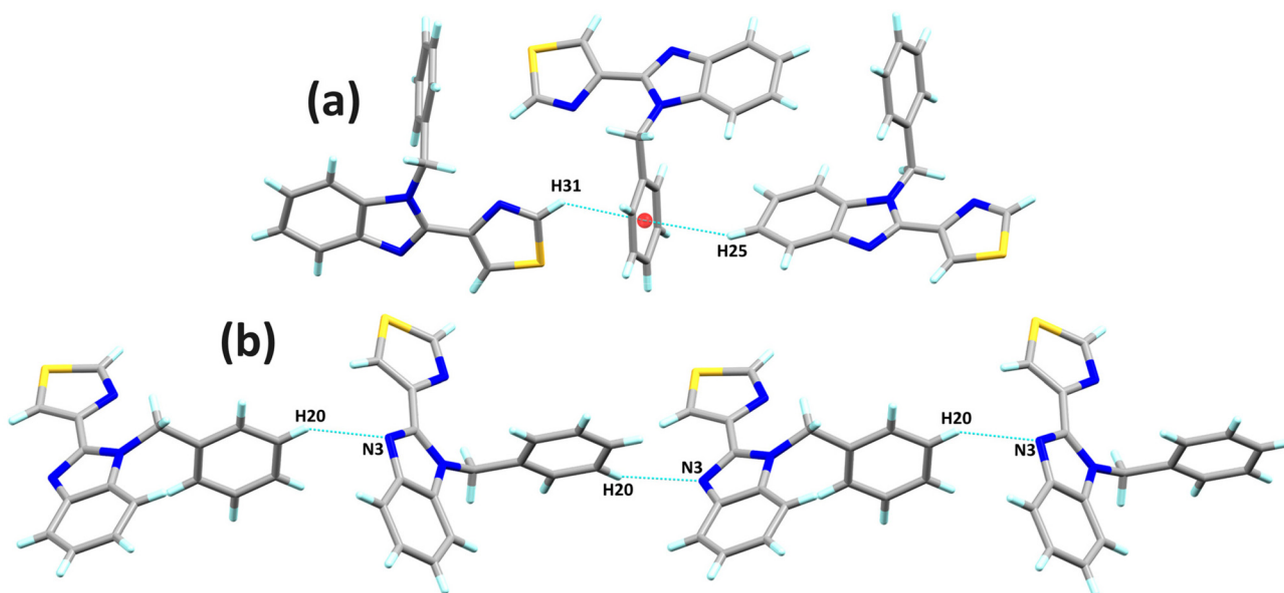
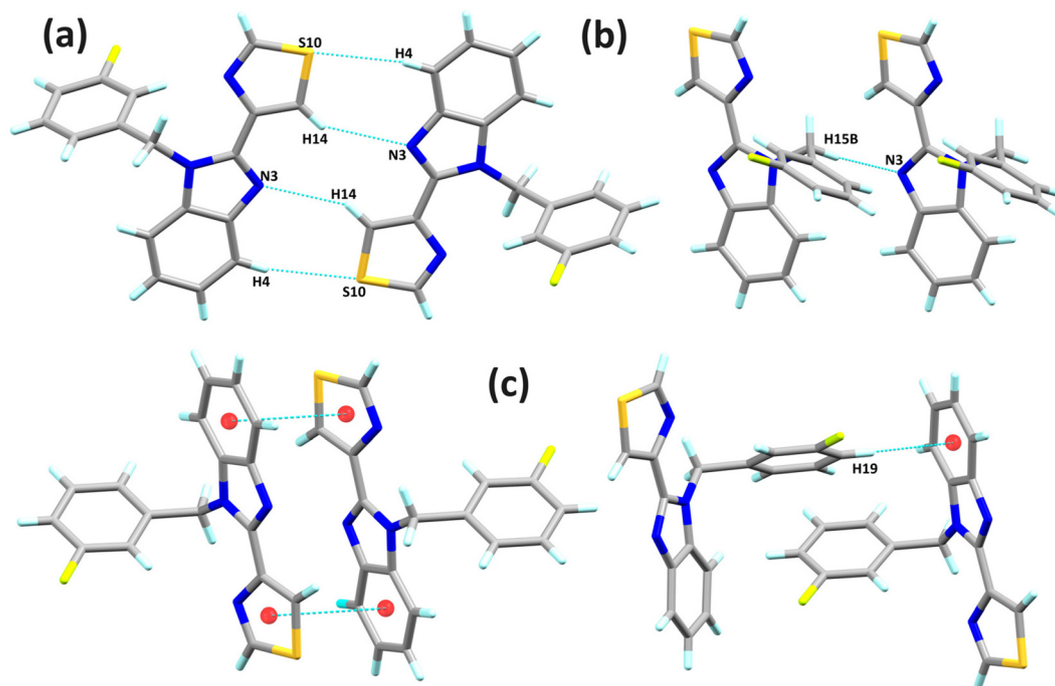


Fig. 2 (a) Lamellar arrangement of L1 parallel to the ac plane, and (b) chain arrangement of L1 generated by the C–H...N interaction.

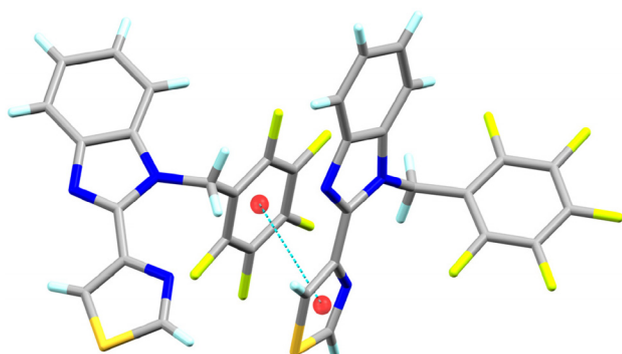
could improve the current treatment of cysticercosis in a murine model with *Taenia crassiceps*.<sup>42</sup> This treatment was dependent on the exposure time and the concentration of

both drugs.<sup>43</sup> In addition, the best time to start treatment was 10 days after infection and at least 20 days of treatment were required.<sup>44</sup> *In vitro* studies of the combination of nitazoxanide





**Fig. 3** (a) Dimeric arrangement of L2 generated by the C-H...N and C-H...S interactions, (b) chain arrangement of L2 generated by the C-H...N interaction, and (c) representation of the  $\pi$ - $\pi$  and C-H... $\pi$  interactions found in L2.



**Fig. 4** Representation of the  $\pi$ - $\pi$  interactions found between the pentafluorobenzyl and thiazole rings in L4.

(NTZ) with ABZ<sup>45</sup> or of NTZ and flubendazole (FLB) could be useful in the treatment of infections caused by *Taenia crassiceps* since they induce a greater metabolic impact on cysticerci compared to the isolated exposure of each of them.<sup>46</sup> Moreover, metabolic acidosis was greater in the group of cysticerci treated with NTZ than in the group treated with ABZ-SO, indicating that this is one of the modes of action used by this drug to induce parasite death.<sup>47</sup> NTZ induced stress in the energetic metabolism in the tricarboxylic acid cycle, in protein catabolism and in the oxidation of fatty acids in *Taenia crassiceps*, so it could be used as an alternative for the treatment of cysticercosis.<sup>48</sup> Additionally, other systems have been studied, such as curcumin oxidation products, which could act as suit-

able inhibitors of thioredoxin-glutathione reductase from *Taenia crassiceps* cysticerci.<sup>49</sup> As well as the ternary system ABZ- $\beta$ CD-PVP (albendazole,  $\beta$ -cyclodextrin and polyvinylpyrrolidone) due to its greater dissolution, high bioavailability and better cysticidal activity with respect to ABZ.<sup>50</sup> Or, treatment with dehydroepiandrosterone (DHEA) which inhibits the establishment, growth and reproduction of *Taenia crassiceps* metacystodes.<sup>51</sup>

From the above, it is evident that most of the studies carried out for the treatment of cysticercosis in murine models with *Taenia crassiceps* have been carried out on drugs with a benzimidazole structure. Furthermore, it is known that evagination is a key indicator of the ability of the scolex to evaginate, a crucial process in the life cycle of *Taenia crassiceps*. Therefore, part of the methodology proposed by Parra-Unda and collaborators was used to evaluate the antiparasitic activity of compounds L1-L5.<sup>52</sup> This test is particularly relevant to this study as it helps understand the compounds' impact on the parasite's ability to develop and survive. In this context, the group incubated with ABZ at a concentration of 500  $\mu$ M showed that after 24 h of incubation, 70% of evaginated cysticerci presented high mobility, but their integrity was reduced from complete to moderate. In the group incubated with TBZ, the percentage of evaginated cysticerci was 100% at the four exposure concentrations, and no reduction in integrity or mobility was observed. The cysticerci were only reduced in size by increasing the incubation time from 24 to 48 h with TBZ. In the five groups of parasites incubated with each of the five compounds, the percentage of evaginated cysticerci was 100%



**Table 2** Effect of ABZ, TBZ and L1–L5 (100, 200, 300 and 500  $\mu\text{M}$ ) on *Taenia crassiceps* cysticerci cultured in RPMI at 24 h

|                   | ABZ                           | TBZ                           | L1                            | L2                            | L3                            | L4                            | L5                            |
|-------------------|-------------------------------|-------------------------------|-------------------------------|-------------------------------|-------------------------------|-------------------------------|-------------------------------|
| Control           | [E] 100%<br>[M] ++++<br>[D] C |
| EtOH/DMSO (0.5%)  | [E] 100%<br>[M] ++++<br>[D] C |
| 100 $\mu\text{M}$ | [E] 85%<br>[M] ++++<br>[D] C  | [E] 100%<br>[M] ++++<br>[D] C | [E] 100%<br>[M] ++++<br>[D] C | [E] 100%<br>[M] ++++<br>[D] C | [E] 100%<br>[M] ++++<br>[D] C | [E] 100%<br>[M] ++++<br>[D] C | [E] 100%<br>[M] ++<br>[D] C   |
| 200 $\mu\text{M}$ | [E] 80%<br>[M] ++++<br>[D] C  | [E] 100%<br>[M] ++++<br>[D] C | [E] 100%<br>[M] ++++<br>[D] C | [E] 100%<br>[M] ++++<br>[D] C | [E] 100%<br>[M] ++++<br>[D] C | [E] 100%<br>[M] ++++<br>[D] C | [E] 100%<br>[M] +<br>[D] C    |
| 300 $\mu\text{M}$ | [E] 75%<br>[M] ++++<br>[D] C  | [E] 100%<br>[M] ++++<br>[D] C | [E] 100%<br>[M] ++++<br>[D] C | [E] 100%<br>[M] ++++<br>[D] C | [E] 100%<br>[M] ++++<br>[D] C | [E] 100%<br>[M] +<br>[D] C    | [E] 100%<br>[M] –<br>[D] C    |
| 500 $\mu\text{M}$ | [E] 70%<br>[M] ++++<br>[D] M  | [E] 100%<br>[M] ++++<br>[D] C | [E] 100%<br>[M] +<br>[D] C    | [E] 100%<br>[M] +<br>[D] C    | [E] 100%<br>[M] +++<br>[D] C  | [E] 100%<br>[M] –<br>[D] C    | [E] 100%<br>[M] –<br>[D] C    |

Viability was determined by: [E] percentage of evaginated cysticerci: number of evaginated parasites/total number of cultured parasites. [M] Parasite mobility: high (++++), medium (+++), moderate (++) and low (+), and null (–). [D] Damage to the cysticerci wall: complete (C), medium (M) and null (N). Cysticerci cultured in RPMI medium were used as a control. Additionally, cysticerci were exposed to 0.5% DMSO for the ABZ assay and 0.5% absolute ethanol for the TBZ and L1–L5 assays.

at any of the concentrations tested. Nonetheless, all compounds reduced the mobility of cysticerci with increasing concentration. L4 and L5 annulled mobility after 24 h of treatment at concentrations of 500  $\mu\text{M}$  and 300  $\mu\text{M}$ , respectively. A special case was the result obtained after 24 hours of incubation with L1 at a concentration of 500  $\mu\text{M}$ , observing 100% of evaginated but wrinkled cysticerci with low mobility. This could be because *N*-benzylation increases the partition coefficient, promoting a species with greater bioavailability but without action on mobility. L4, with the highest amount of fluorine substituents in the *N*-benzyl fragment, presented greater activity at a lower concentration compared to L1–L3 but lower compared to L5. The steric volume and electron density of fluorine atoms in L4 could be a critical factor in the interaction with phenylalanine of parasitic  $\beta$ -tubulin. L5, with the trifluoromethyl group in the *para* position of the *N*-benzyl ring, presented a greater action at a concentration comparable to that of ABZ, which could be directly related to the cancellation of the parasites' mobility, preventing their transfer to the hatching zone. L5 is an excellent candidate for future studies in the treatment of cysticercosis. This compound presents the trifluoromethyl group in its structure, a very important functional group in recent studies of the possible drugs RCB15 and RCB20, Table 2.

### Ethics approval

All protocols were carried out in the animal facilities of the Facultad de Medicina Universidad Nacional Autónoma de México, under controlled conditions of temperature (22 °C), a relative humidity of 50–60% and 12 h dark/light cycles, in strict accordance with the Official Mexican Norm for the Production, Care, and Use of Laboratory Animals (NOM-062-ZOO-1999) and the Guide for the Care and Use of Laboratory

Animals of the National Institutes of Health, U.S.A. The project was also approved by the Internal Committee for the Care and Use of Laboratory Animals (CICUAL-020-CIC-2018) de la Facultad de Medicina, UNAM, Mexico.

## Conclusions

Five compounds derived from TBZ with an *N*-benzyl arm functionalised with fluoride substituents were synthesised and characterised. Remarkably, four of the compounds, excluding L1, have not been previously reported, adding a significant element of novelty to our research, as confirmed by our query in CAS Scifinder<sup>®</sup>. Crystals of L1, L2 and L4 suitable for characterisation by SC-XRD were obtained. The supramolecular arrangements of the three structures are mainly supported by C–H...N3 intermolecular interactions. In addition, other weak interactions are reported, such as C–H...S10, C–H... $\pi$  and  $\pi$ ... $\pi$ . All compounds showed activity in the mobility of *Taenia crassiceps* cysticerci, especially L4 and L5. The effect of ABZ on mobility, integrity and evagination was observed at different concentrations. ABZ is the drug of choice to treat neurocysticercosis, acting selectively on the cytoplasmic microtubules of the intestinal cells of the parasite. TBZ inhibits the fumarate reductase enzyme of some helminths other than *Taenia solium* and *Taenia crassiceps*. Thus, this investigation represents the first study against cysticercosis in a murine model with *Taenia crassiceps* using novel, non-commercial molecules.

## Author contributions

M. I. Rodríguez-Mora: investigation, methodology and conceptualization. R. Colorado-Peralta: writing – original draft,



visualization and formal analysis. V. Reyes-Márquez: writing – original draft, visualization and formal analysis. M. A. García-Eleno: investigation, supervision, software and validation. E. Cuevas-Yáñez: investigation, supervision, software and validation. J. R. Parra-Unda: investigation, methodology, resources and data curation. A. Landa: investigation, methodology, resources and data curation. D. Morales-Morales: conceptualization, resources, funding acquisition, project administration, and writing – review and editing.

## Data availability

The data supporting this article have been included as part of the ESI.†

## Conflicts of interest

There are no conflicts to declare.

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