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Effect-directed analysis and chemical identification of agonists of peroxisome proliferator-activated receptors in white button mushroom†

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Obesity has a serious effect on human health. It relates to metabolic syndrome, including the associated disorders such as type 2 diabetes, heart disease, stroke and hyperemia. The peroxisome proliferator-activated receptors (PPARs) are important receptors to control fat metabolism in the human body. Because of the safety concerns of synthetic drugs targeting PPARs, ligands from natural sources have drawn interest. Earlier, we have found high PPAR activities in extracts from *Agaricus bisporus* (white button mushroom, WBM). WBM contains a wide range of candidate compounds which could be agonists of PPARs. To identify which compounds are responsible for PPAR activation by WBM extracts, we used fractionation coupled to effect-directed analysis with reporter gene assays specific for all three PPARs for purification and LC/MS-TOF and NMR for compound identification in purified active fractions. Surprisingly, we identified the relatively common dietary fatty acid, linoleic acid, as the main ligand of PPARs in WBM. Possibly, the relatively low levels of linoleic acid in WBM are sufficient and instrumental in inducing its anti-obesogenic effects, avoiding high energy intake and negative health effects associated with high levels of linoleic acid consumption. However, it could not be excluded that a minor relatively potent compound contributes towards PPAR activation, while the anti-obesity effects of WBM may be further enhanced by receptor expression modulating compounds or compounds with completely PPAR unrelated modes of action.

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Introduction

Obesity has a significant impact on human health, leading to an increased incidence of metabolic syndrome, which is characterized by central obesity, high blood pressure, high blood sugar, high level of triglycerides, and low serum high-density lipoprotein (HDL). This can lead to type 2 diabetes,

heart disease and hyperemia.^{1–3} Thus, there is a great demand for safe treatments of these disorders. Several drugs are used to treat obesity-related disorders. These include fibrates to treat hyperlipidemia and thiazolidinediones (TZD) to treat type 2 diabetes. Remarkably, these drugs were found to be ligands for peroxisome proliferator-activated receptors (PPARs). PPARs are a subgroup of nuclear hormone receptors and are essential in the regulation of fat storage and metabolism.^{4,5} There are three isoforms of PPARs, *i.e.*, PPAR α , PPAR γ , and PPAR δ , with partly overlapping and specific functions, and different tissue distribution. PPAR α is a key player in lipid metabolism, while PPAR γ has a major role in stimulating adipogenesis and improving insulin sensitivity. Fibrates are ligands for PPAR α , while TZD targets PPAR γ .⁵ PPAR δ also is important in fat metabolism, generating heat when uncoupled to energy production, playing a role in the switch from glucose to fat burning.^{4,5} Unfortunately there are serious side effects related to the use of TZDs such as rosiglitazone, while so far attempts have failed to generate safe drugs for the most interesting receptor in this context, PPAR δ .^{6,7} There is a great need to identify alternative PPAR ligands, which could include non-

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drug related approaches. It should be noted that there is a possibility that bioactive compounds activating the PPARs would give the same side effects as the current pharmaceuticals, if these side effects were inherent to PPAR activation. This indeed could be the case, and there are concerns and controversies regarding the effects of PPAR δ activation on cancer progression.⁸ This, however, may be caused by the complex nature of cancer progression and the many possible points of interaction, together with the differences in the animal and cellular models used.⁸ To avoid misinterpretation of these data, the best possible test should be carried out in humans. Since this is not feasible, an alternative is the identification of active molecules in a food item with a long history of safe use, as has been done in the current study.

As such, life-style changes, including more healthy diets and dietary supplements, are increasingly seen as efficient alternatives to pharmacological treatment.⁹ Edible mushrooms have been reported to be effective in combating metabolic disorders including reduction of hyperlipidemia, hypertension, and hyperglycemia, and are a possible resource to isolate their functional ingredients avoiding the side effects of synthetic drugs.^{10,11} Humans have used edible mushrooms, including *Agaricus bisporus*, *Pleurotus ostreatus*, and *Lentinula edodes*, for hundreds of years because of their health benefits.^{12–15} White button mushroom (WBM; *Agaricus bisporus*) is the most produced mushroom worldwide, and could be an interesting source of anti-obesogenic molecules.^{16,17} It has been shown in rats that a WBM supplement to a high-fat diet resulted in a significant decrease in body weight after 72 days, and inhibited liver accumulation of fat.¹⁸ In addition, WBM powder fed to diabetic male Sprague-Dawley rats caused a significant reduction of plasma glucose and triglyceride levels after three weeks. Besides, hypercholesterolemic rats fed with WBM powder showed a significant decrease in plasma total cholesterol and low-density lipoprotein.¹⁹ Moreover in a randomized clinical trial in humans, WBM consumption resulted in clear positive effects on body weight and health parameters such as central obesity, blood pressure, and lipid profiles.²⁰

WBM contains a wide range of bioactive compounds, including phenolic compounds, unsaturated and saturated fatty acids, and β -glucans.¹⁶ It is unclear however, which components may be responsible for the possible anti-obesogenic effects observed in humans.²⁰ As always, the cause and significance of these health effects are extremely difficult to substantiate in a background of variable and complex food intake. A wide range of these natural compounds, including fatty acids, polyphenols, and flavonoids, have been shown to be ligands of PPARs, thus making these receptors interesting targets to discover natural compounds that are active in the prevention or treatment of metabolic diseases.^{7,21} To this end, we recently developed a specific panel of human cell-based CALUX® reporter gene assays for all three full-length PPARs and found high PPAR activities in fungi.¹⁷ Notably, high levels of PPAR δ activation were found in edible mushroom including the common WBM.¹⁷ Here, we used effect-directed analysis and chemical identification using liquid chromatography

coupled time-of-flight mass spectrometry (LC/MS-TOF) and nuclear magnetic resonance spectroscopy (NMR) to elucidate the identity of the PPAR interacting ligands from WBM.

Materials and methods

WBM powder preparation

WBM was purchased from a local supermarket in The Netherlands. Two kilograms of fresh WBM was cut into small pieces and dehydrated by using a freeze drier. Then, the dehydrated WBM was ground to a powder using an A11 Basic analytical mill (IKA, Staufen, Germany). This standardized batch of WBM powder, used in all experiments, was stored at -80 °C until further use.

Chemicals

Rosiglitazone ($\geq 98\%$, CAS no. 122320-73-4) was purchased from Bio-connect BV (Huissen, The Netherlands). L-0165.041 ($\geq 98\%$, CAS no. 79558-09-1) and GW7647 ($\geq 98\%$, CAS no. 25129-71-3) were purchased from Sigma-Aldrich (Zwijndrecht, The Netherlands). Methanol (absolute, CAS no. 67-56-1) and acetonitrile (99.9%, CAS no. 75-05-8) were purchased from Biosolve BV (Valkenswaard, The Netherlands). Ethanol ($\geq 99.5\%$, CAS no. 64-17-5) was obtained from Thermo Fisher Scientific BV (Breda, The Netherlands). Dimethylsulfoxide (DMSO, 99.9%, CAS no. 67-68-5) was obtained from Thermo Fisher Scientific BV (Breda, The Netherlands). Linoleic acid (LA) ($\geq 99\%$, CAS no. 60-33-3), conjugated linoleic acid (CLA) 9-*cis*, 11-*trans* ($\geq 96\%$, CAS no. 2540-56-9), and CLA 10-*trans*, 12-*cis* ($\geq 96\%$, CAS no. 2420-56-6) were purchased from Sigma-Aldrich (Zwijndrecht, The Netherlands). Water used for extraction and fractionation was purified using a PureLab flex system (ELGA Labwater, High Wycombe, UK).

Preparation of WBM extracts

The extraction procedure was a modification of the method developed by Gijsbers *et al.*;²² in brief: 3 mL of 0.1 M sodium acetate (pH 4.8) and 1 mL of viscozyme L (Sigma-Aldrich, St Louis, MO, USA) were added to 1 g of the dried sample material, and the mixture was incubated at 37 °C for 2 hours. Then, 30 mL of ethanol was added, and the sample mixture was transferred to an ultrasonic bath for 10 minutes and subsequently centrifuged at 3000g for 10 minutes. The supernatant was collected and dried in a GeneVac Rocket evaporation system (Genevac Limited, Ipswich, UK). The WBM dried extract was re-dissolved in 2 mL methanol and filtered. The mixture was kept at -20 °C until further experiments.

Cell culture

CALUX cells based on human U2OS osteosarcoma cells (American Type Culture Collection [ATCC], Manassas, USA), expressed a full length PPAR α , δ or γ receptor construct, respectively, and a luciferase reporter gene.¹⁷ In addition, to assess non-specific luciferase modulation, *e.g.* due to cytotoxicity, the Cytotox CALUX cell line was used, consisting of



U2-OS cells constitutively expressing the luciferase gene.²³ CALUX cells were cultured in a DMEM/F12 glutamax medium supplemented with 7.5% fetal calf serum, non-essential amino acids, and penicillin/streptomycin (all from Invitrogen, Breda, The Netherlands) at the final concentrations of 10 U mL⁻¹ and 10 µg mL⁻¹, respectively, as described before.²⁴ Once per week, G418 (Duchefa Biochemie, Haarlem, The Netherlands) was added to the culture medium at a nominal concentration of 200 µg mL⁻¹.

CALUX® reporter gene assays

The PPAR activities of sample extracts and fractions were tested by measuring luciferase activity in the PPAR CALUX reporter cells. In parallel, the Cytotox CALUX was used to assess assay interferences. Samples that led to 20% or more signal reduction were excluded from the evaluation.²³ PPAR CALUX reporter gene assays were performed as described before.¹⁷ In short, PPAR CALUX cells were seeded in 96-well plates (Greiner-Bio-One, Alphen aan den Rijn, The Netherlands) at a density of 10 000 cells per well in a 100 µL assay medium (DMEM/F12 without phenol red, Thermo Fisher Scientific BV, Breda, The Netherlands). After 24 hours, when the cells formed a monolayer, 100 µL of fresh assay medium supplemented with the sample extract or WBM fractions were added to the wells. When testing the sample, the percentage of DMSO in the exposure medium was kept at 0.1%. On each plate, a dilution series of reference PPAR agonists (rosiglitazone, L-0165.041 and GW7647) was included as a reference curve. After 24 hours of exposure, the light production as relative light units (RLU) of luciferase activity was measured using a TriStar LB 941 luminometer (Berthold Technologies, Bad Wildbad, Germany).

PPAR CALUX activity in the sample extracts was expressed as a relative induction to the maximum response from the reference compounds. These results were interpolated in the PPAR CALUX bioassay specific calibration curves using the statistical software package GraphPad Prism V5.03 (non-linear regression, variable slope, 4 parameters, robust fit), after which the bioactivity was quantified. For each of the sample extracts, the lowest concentration of the sample extract present in the CALUX exposure medium (µg sample per L medium) causing a relative induction of 10% (PC10) was determined as the lowest observed effect concentration (LOEC).

Sample fractionation and effect-directed analysis

WBM fractionation was carried out using three LC columns (biphenyl semi-prep column, F5 column, and C18 column; Fig. 1), using a HPLC Agilent 1260 Infinity system (Agilent Technologies Netherlands BV, Amstelveen, The Netherlands). First, the WBM extract was fractionated using a biphenyl semi-prep column (Kinetex 5 µM biphenyl, 250 × 10 mm; Phenomenex, Torrance, USA). A gradient solvent system using methanol:water was used from 20% to 95% for 27 minutes, with a flow rate of 3.8 mL per minute. Subsequent fractions were collected using a Waters fraction collector III (Waters Corporation, Milford, USA) and evaluated for PPAR activities

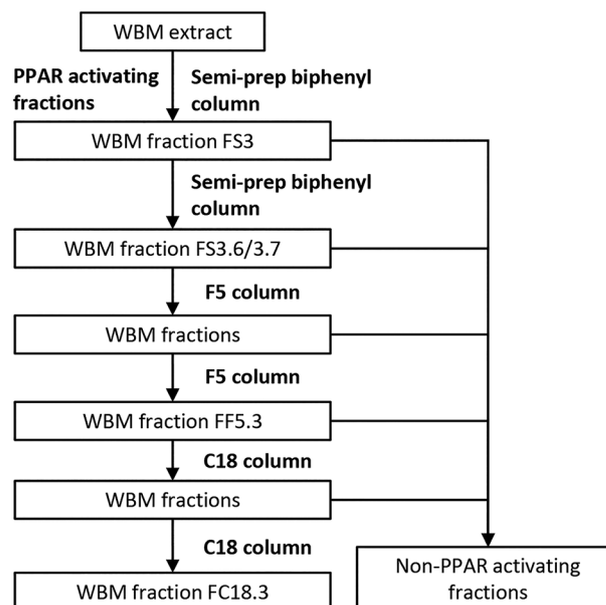


Fig. 1 Flowchart of fractionation of the WBM extract using a series of LC columns (semi-prep biphenyl column, F5 column, and C18 column). All PPAR activities of WBM fractions were determined by PPAR assays.

by CALUX assays. WBM fraction 3 (FS3), which was the highest in PPAR activity, was further sub-fractionated using the biphenyl semi-prep column using smaller, one-minute intervals of collection. WBM active fraction FS3.6 and FS3.7 were separated further on an analytical F5 column (Kinetex 1.7 µM F5, 100 × 2.1 mm; Phenomenex, Torrance, USA), using a solvent gradient of methanol:water from 20% to 100% in 31 minutes, with a flow rate of 0.25 mL per minute, and fractions were tested for bioactivities using PPAR CALUX assays. The WBM active fraction from the F5 column was further sub-fractionated on an F5 column. In the last step of WBM ligand purification, the active fraction collected from the F5 (FF5.3) column was purified by using a C18 column (Kinetex 2.6 µM UXB-C18, 75 × 2.1 mm; Phenomenex, Torrance, USA). A solvent gradient was applied using acetonitrile (0.1% formic acid):water (0.1% formic acid) from 70% to 90% in 10 minutes, with a flow rate of 0.25 mL per minute. WBM active fraction from the C18 column was fractionated a second time using the C18 column and the final highly purified PPAR active fraction FC18.3 obtained was dissolved in acetonitrile and kept at -20 °C.

LC/MS-TOF analysis and data processing

The WBM active fractions were analyzed by LC/MS-TOF and data were processed by non-target analysis as described before;²⁵ in brief: the fractions were analyzed using a Nexera UHPLC system (Shimadzu, Den Bosch, The Netherlands), coupled to a maXis 4G high-resolution quadrupole time-of-flight HRMS (UHPLC-Q-TOF-MS/MS), equipped with a HD collision cell (Bruker, Leiderdorp, The Netherlands). Liquid chromatography (LC) analysis was performed according to the aforementioned programs for Kinetex F5 column and Kinetex



C18 column. 2 mM sodium formate in 1 : 1 methanol : water solution was used for the automatic internal calibration of the system. Positive and negative ESI were used to acquire MS- and MS/MS data in the range of 80–1200 *m/z*. A UV chromatogram (210 nm) was simultaneously recorded to detect the active peak of the samples. The peak and retention times in the UV chromatogram were compared to those found before in the F5 column and C18 column. Non-target analysis (NTA) was performed for processing the LC/MS-TOF data using patRoom, which is an open-source platform that harmonizes various commonly used software tools employed in NTA.²⁶ The NTA workflow has five steps, as follows (see ESI 1†): at first, the LC/MS-TOF raw data were converted to the open mzML format.²⁷ Secondly, feature data were extracted and grouped by OpenMS algorithms.²⁸ Third, the features were filtered by different constraints (see ESI 1†). Fourth, the filtered features were automatically annotated by extracting mass spectral data using the mzR algorithm.²⁹ Formulae were automatically calculated for all the features with both the GenForm and SIRIUS algorithms.^{30–34} Compound structure annotation was performed with SIRIUS and MetFrag,³⁵ both using the PubChem library.³⁶ Only the top 25 ranked candidates with an explained chemical formula were kept. The last step, “componentization”, was performed to automatically detect the features that are isotopes or MS adducts (using RAMClustR as an algorithm).³⁷

NMR analysis

¹H and ¹³C{¹H} NMR spectra of the dilute solution in the CDCl₃ solvent were recorded on a Bruker Avance Neo-500 MHz instrument and calibrated based on residual undeuterated solvent signals as an internal standard. ¹³C{¹H} NMR spectra of 16 384 transients were measured by broadband decoupling and at a relaxation delay of 2.0 s. 2D spectra were recorded with 2K/256 data points. The 2D homonuclear shift correlation was established using gradient pulses for the selection. The HSQC spectrum probed the 2D ¹H-/¹³C correlation *via* double

inept transfer utilizing sensitivity improvement, phase sensitivity using echo/antiecho-TPPI gradient selection with decoupling during acquisition and multiplicity editing during the selection step.^{38–42}

Data analysis

PPAR CALUX activities and associated PPAR biological equivalents (BEQs) were expressed as mean ± standard deviation (SD). One-way ANOVA was employed to determine statistical significance. Statistical significance was set at *P* < 0.05.

Results and discussion

Bioassay-guided fractionation of bio-active compounds from WBM

In order to purify and identify the compounds responsible for PPAR activation in WBM extracts,¹⁷ we set out to separate and fractionate a standardized sample. Through linkage of fractionated samples with bioactivity, we purified the compound(s) of interest stepwise. In this study, we used HPLC to fractionate the WBM extracts and determined all three PPAR activities using CALUX bio-assays specific for PPAR α , PPAR δ and PPAR γ . First, a standardized WBM extract was prepared to be used in all subsequent experiments. It was extracted and tested for the presence of PPAR ligands using the CALUX assays. The activity profile was similar to the one noted before.¹⁷ Again, a high concentration of PPAR agonist activity was measured confirming the quality of the WBM batch to be suitable for further experimentation (Table 1).

Note that the activity of the PPAR δ extract is relatively high as compared to the other PPARs when expressed as biological equivalents (BEQs). This is also because the reference ligand is of lower potency than the available ones for PPAR α and PPAR γ , which stresses the fact that ligands for PPAR δ are less well developed and that there is a great need to search for alternatives,⁴³ as has been done in the current study. The PPAR activities (as expressed as BEQs) tended to be higher in the purified

Table 1 PPAR α , PPAR δ and PPAR γ activity in a standardized WBM extract and purified fractions

	PPAR α			PPAR δ			PPAR γ		
	LOEC at PC10 ($\mu\text{g L}^{-1}$ medium)	BEQ (ng GW7647-eq. per g dry WBM)	Maximum activity of PPAR α samples (%)	LOEC at PC10 ($\mu\text{g L}^{-1}$ medium)	BEQ (ng L-165.041 eq. per g)	Maximum activity of PPAR δ samples (%)	LOEC at PC10 ($\mu\text{g L}^{-1}$ medium)	BEQ (ng rosiglitazone eq. per g dry WBM)	Maximum activity of PPAR γ samples (%)
WBM extract	190 546	154 ± 8	40.1 ± 3.3	—	1598 ± 203	8.9 ± 0.9	—	127 ± 11	2.8 ± 0.5
FS3	1 181 436	459 ± 54*	57.9 ± 7.6*	4 478 967	3800 ± 407*	17.0 ± 0.7*	3 517 214	660 ± 56*	16.9 ± 1.7*
FS3.6	329 864	600 ± 48*	67.6 ± 7.5*	1 249 930	3400 ± 126*	19.9 ± 1.0*	2 797 986	360 ± 30*	26.1 ± 1.5*
FS3.7	524 406	620 ± 105*	49.0 ± 7.8	3 138 435	1500 ± 101	18.6 ± 1.3*	6 660 172	170 ± 3	11.6 ± 0.3*
FF5.3	154 387	300 ± 31*	61.0 ± 7.3*	901 961	6000 ± 354*	23.7 ± 1.2*	1 659 570	230 ± 11*	11.0 ± 0.5*
FC18.3	155 597	340 ± 12*	75.4 ± 3.5*	1 218 990	6200 ± 403*	45.4 ± 6.2*	6 295 062	260 ± 30*	12.4 ± 1.8*

The lowest observed effect concentration (LOEC) of samples were expressed as microgram dry WBM per liter medium. PPAR BEQs of samples were expressed relative to GW7647 (PPAR α agonist reference compound), L-165.041 (PPAR δ agonist reference compound), and rosiglitazone (PPAR γ agonist reference compound), respectively. Luciferase activity of samples was expressed as a percentage of the maximum PPAR reference compound response. Data are corrected for solvent control luciferase values and are expressed as mean ± SD (*n* = 3). *: Significant difference compared with the WBM extract, *P* < 0.05.



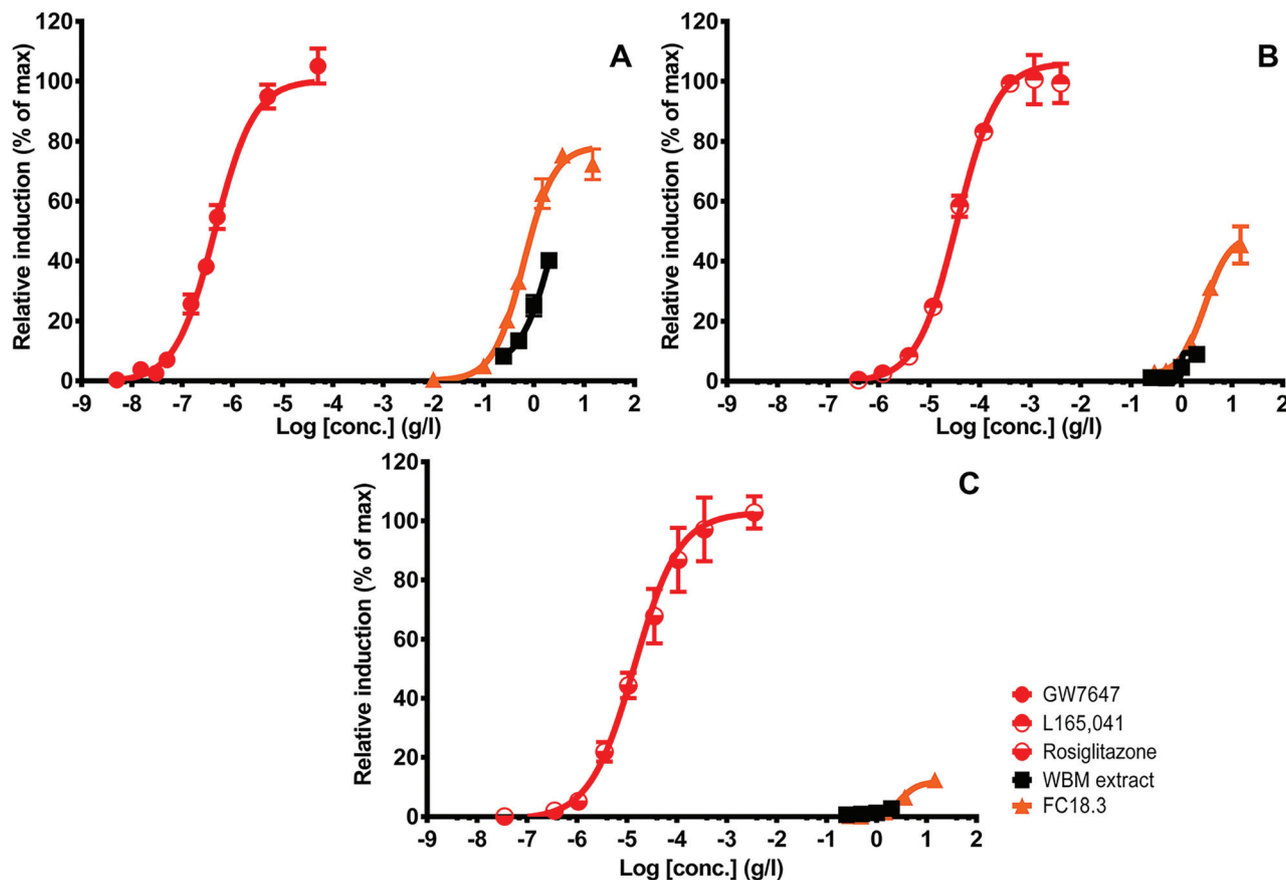


Fig. 2 Concentration-dependent increase in PPAR α , PPAR δ and PPAR γ activity upon exposure to crude WBM extract and purified WBM FC18.3. PPAR α luciferase activity is expressed as a percentage of the maximum GW7647 (A) response. PPAR δ luciferase activity is expressed as a percentage of the maximum L-165,041 (B) response. PPAR γ luciferase activity is expressed as a percentage of the maximum rosiglitazone (C) response. All WBM concentration points passing the threshold of cytotoxicity to the CALUX cells were excluded from the graph, resulting in shorter dose–response curves in the crude fraction. Data are corrected for solvent control luciferase values and are expressed as mean \pm SD in triplicate ($n = 3$).

fractions compared to the original extract. In general, concentration response curves were more complete and cytotoxicity was reduced (Fig. 2; note that fewer data points were excluded because of cytotoxicity in the purified fraction).

First, the WBM extract was fractionated using the semi-prep biphenyl column and five fractions (FS1 to FS5) were collected (Fig. 3). We chose this column since this seemed to be the most suitable one to separate the various molecules known to be produced by WBM, and since this has been shown to interact with PPARs like polyphenols and lipids.¹⁶ Among the five fractions obtained, fraction 3 (FS3) contained the highest PPAR BEQ levels (Fig. 3). The PPAR BEQ values were significantly higher than that of the original unfractionated WBM extract (Table 1), with PPAR α levels 3-fold, PPAR δ levels 2-fold, and PPAR γ levels 5-fold higher, suggesting that all relevant molecules are present in this fraction. The higher activity may be explained by a loss of interfering molecules.

To get a more purified fraction, WBM FS3 was further sub-fractionated using smaller, one-minute intervals of collection. The highest PPAR activity was found in two fractions of one minute, starting from 19 minutes to 21 minutes (FS3.6–FS3.7;

Fig. 4), with the highest activity in FS3.6. The chromatogram of the sub-fractionation showed one peak co-eluting with the main activity in FS3, as present in FS3.6 and FS3.7. Therefore, FS3.6 and FS3.7 were pooled to one fraction for further separation.

For further purification, we ran the WBM fraction on an analytical F5 column, which provides a different interaction to the semi-prep biphenyl column, resulting in a high efficiency in separation and purity of the identified active fraction. The WBM fractionation process on the F5 column also proceeded, at first, in a small number of relatively large fraction volumes pooled, followed by a more diligent smaller step fractionation. The highest PPAR activity of the biphenyl fraction FS3.6/3.7 separated on the F5 column was in fraction FF5.3 (Fig. 5). At the end of this procedure, the purity of the WBM fraction collected from the F5 column was considered enough for structural elucidation by LC/MS-TOF.

LC/MS-TOF identification of PPAR agonists in the purified WBM extract

From the LC/MS-TOF analysis of fraction FF5.3, over 30 features were detected (see ESI 2, Table 2-1; 2-2[†]). Most features



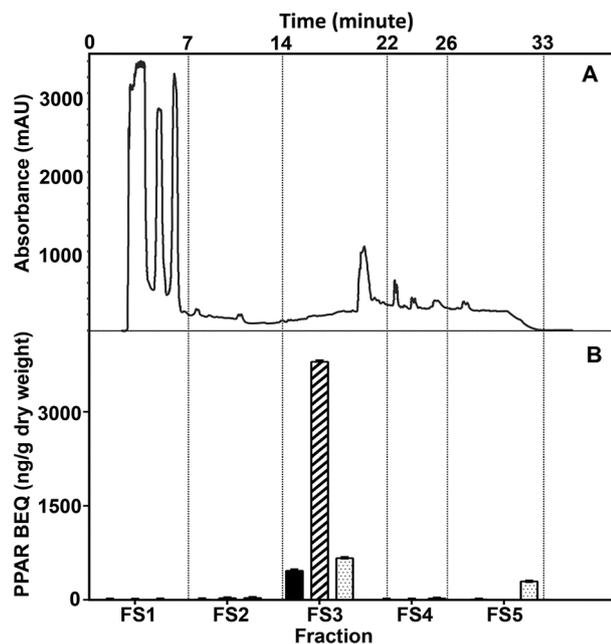


Fig. 3 Chromatogram of WBM extract separated on a semi-prep biphenyl column (A) and determination of PPAR BEQs in five fractions (FS1–FS5; B). The PPAR α BEQ values of fractions were expressed relative to GW7647 (black bars). The PPAR δ BEQ values of fractions were calculated relative to L-165.041 (cross-hatched bars). The PPAR γ BEQ values of fractions were expressed relative to rosiglitazone (dotted bars). The data are expressed as mean \pm SD in triplicate ($n = 3$).

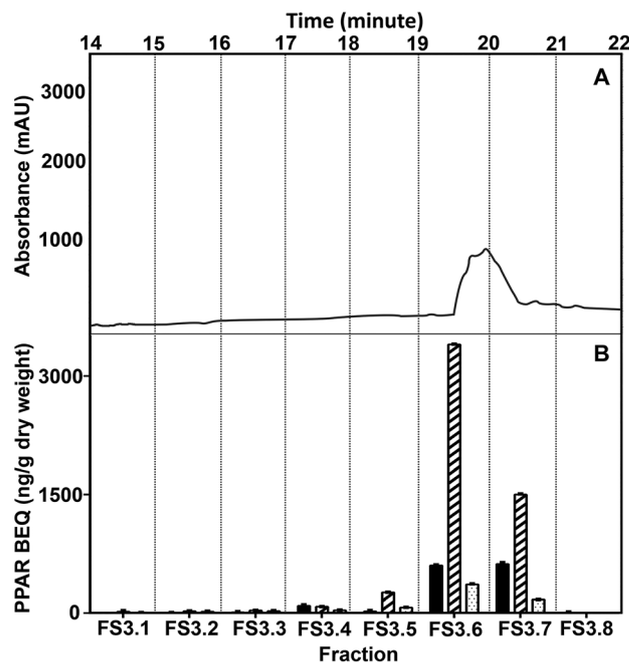


Fig. 4 Chromatogram of sub-fractionation of the active fraction (FS3) from the WBM extract separated using a semi-prep biphenyl column (A) and determination of PPAR BEQ in eight fractions (FS3.1–FS3.8; B). The PPAR α BEQ values of fractions were expressed relative to GW7647 (black bars). The PPAR δ BEQ values of fractions were calculated relative to L-165.041 (cross-hatched bars). The PPAR γ BEQ values of fractions were expressed relative to rosiglitazone (dotted bars). The data are expressed as mean \pm SD in triplicate ($n = 3$).

(19) were detected with negative ionization, which suggests a predominant presence of acidic compounds. Due to the large number of features, we did not attempt to perform full structural identification for this fraction. Regardless, basic identification of each feature was performed by automatic calculation of chemical formulae. The high hydrogen to carbon ratios of most formulae suggested primarily aliphatic compounds, and from these most formulae only contained the elements C, H, and O, which suggests fatty acid-like structures.

To further reduce the complexity of the FF5.3 fraction, subsequent fractionation was performed with a C18 column since this stationary phase is highly suitable for the separation of fatty acid-like compounds.⁴⁴ FC18.3 showed a single peak, which correlated with high BEQ of PPAR α and PPAR δ activity (Fig. 6). This further purified fraction WBM FC18.3 was analyzed by LC/MS-TOF, and only a total of 7 features were detected (see ESI 2, Table 2-3[†]). With positive ionization, the four detected features were of similar intensity; however, negative ionization revealed a single most abundant feature (FC18.3-neg-1, 88% relative intensity). Chemical annotation results from the applied workflow suggested linoleic acid (LA) as the top candidate for the most abundant features detected in both positive and negative ionization; hence, we pursued our studies with this compound.

Fig. 7 shows that the retention times of the FC18.3 active fraction compared to those of LA or its major isoforms CLA 9-

cis, 11-*trans* and CLA 10-*trans*, 12-*cis* are close for the latter two and are identical for LA itself.

Since LA and isoforms of LA have been reported to interact with PPARs, these are potential candidates that contribute to WBM-derived activity.^{45,46} Several studies provide evidence of health benefits of LA or its conjugated isoforms (CLAs), including its anti-obesity potential. The most active and well-studied isoforms of LA are CLA 9 *cis*-11 *trans*, and CLA 10 *trans*-12 *cis*, showing a variety of relevant beneficial biological effects, including reduction of type 2 diabetes and body weight.^{47,48} We next studied if these compounds could be related to the molecules in the WBM fraction responsible for PPAR activation. We tested them in the panel of PPAR assays in comparison with the activity of WBM FC18.3, assuming that all activity was LA-derived (Fig. 8). The concentration response curves of WBM FC18.3, expressed as LA equivalents, were quite similar to those of LA and CLA 9-*cis*, 11-*trans*. On the other hand, the PPAR activity response curve of CLA 10-*trans*, 12-*cis* was clearly different compared to the other samples (Fig. 8). Also, when the lowest observed effect concentrations (LOECs) were used, CLA 10-*trans*, 12-*cis* was clearly different from the rest, and this isoform seems not to be responsible for PPAR activation by WBM extracts (Table 2).

Fig. 8 shows that there was a difference in shape between the reference curve and the tested samples that could be



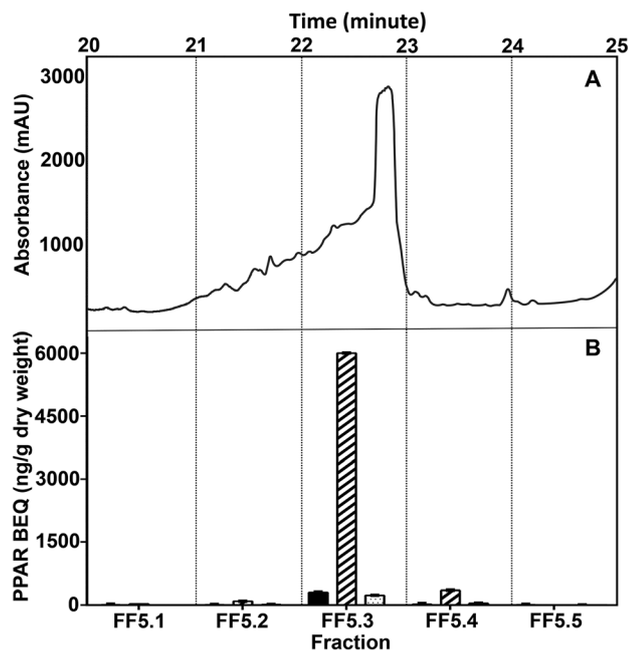


Fig. 5 Chromatogram of WBM fraction FS3.6/3.7 separated on an F5 column (A) and determination of PPAR BEQs in five fractions (FF5.1–FF5.5; B). The PPAR α BEQ values of fractions were expressed relative to GW7647 (black bars). The PPAR δ BEQ values of fractions were calculated relative to L-165.041 (cross-hatched bars). The PPAR γ BEQ values of fractions were expressed relative to rosiglitazone (dotted bars). The data are expressed as mean \pm SD in triplicate ($n = 3$).

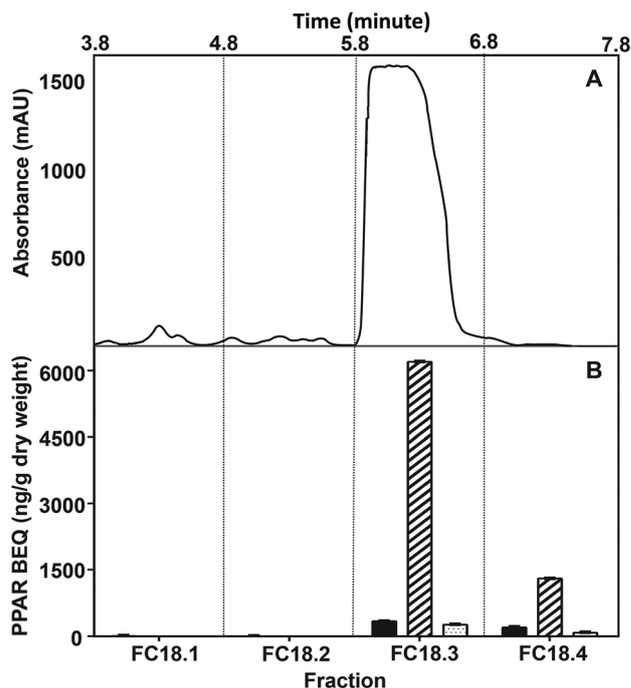


Fig. 6 Chromatogram of WBM FF5.3 fraction separated on a C18 column (A) and determination of PPAR BEQs in four fractions (FC18.1–FC18.4; B). The PPAR α BEQ values of fractions were expressed relative to GW7647 (black bars). The PPAR δ BEQ values of fractions were calculated relative to L-165.041 (cross-hatched bars). The PPAR γ BEQ values of fractions were expressed relative to rosiglitazone (dotted bars). The data are expressed as mean \pm SD in triplicate ($n = 3$).

caused by the fact that the LA-type molecules act at high dosage and never reach their maximum, since at these high dosages cytotoxicity comes into play. Alternatively, the inherent efficacy for receptor activation could be different. In general, the lower end of the concentration response curves of the purest extract (FC18.3) and LA is quite similar, and the curves deviate at the higher dosages (Fig. 8), which suggests that this may be caused by impurities that differ between these samples. When comparing the maximal activity between WBM FC18.3 and the different compounds, again, the LA and CLA 9-*cis*, 11-*trans* profiles were again quite similar, while CLA 10-*trans*, 12-*cis* was clearly different. An exception seems to be the relatively strong effect of WBM FC18.3 towards PPAR δ (Fig. 8 and Table 2).

In conclusion, the LC/MS-TOF results and the candidate molecule testing could not reveal the exact LA isoform and the tested forms are only a minority of the many possible linoleic acid isoforms, all having different biological activities.⁴⁹ For this reason, further characterization with NMR was pursued to determine the isoform(s) responsible for PPAR activation more precisely.

NMR characterization of PPAR agonists from WBM

The ^1H NMR spectrum of the CDCl_3 diluted sample shows an almost pure compound with the characteristic features of an unsaturated olefinic acid, *i.e.* a broad signal of the carboxylic acid proton at a low field and four olefinic protons indicating two double bonds were found to be present. The characteristic

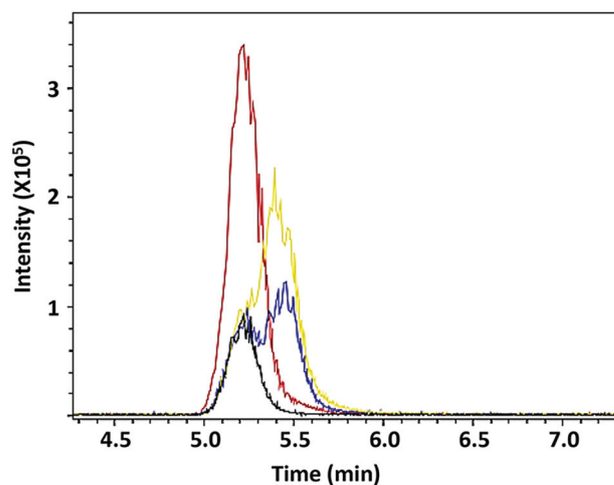


Fig. 7 Overlaid extracted ion chromatograms (positive mode) for sample FC18.3 (black) spiked with linoleic acid (LA, red), CLA 10-*trans*, 12-*cis* (yellow) and CLA 9-*cis*, 11-*trans* (blue).

methylene signal at 2.80 ppm shows only correlation to these olefinic protons, proving the olefinic acid to be non-conjugated. The NMR spectroscopic data (^1H NMR (500 MHz, CDCl_3 , 298 K) δ 10.73 (broad, 1H, HOOC), 5.38 (m, 4H, HC=CH), 2.80 (dd, $J = 6.6$ Hz, 2H, =CH-CH₂-CH=), 2.37 (t, J



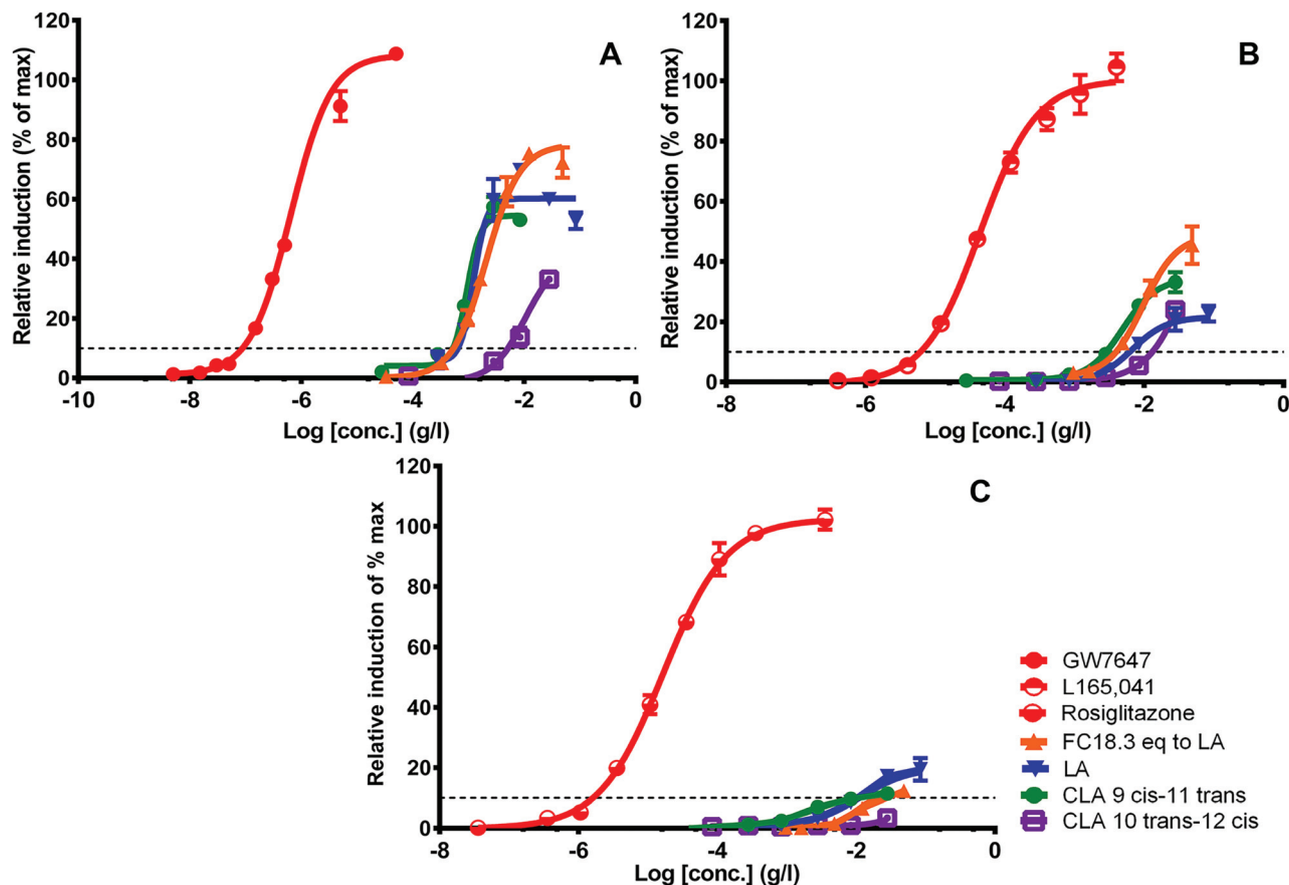


Fig. 8 Concentration-dependent increase in PPAR α , PPAR δ and PPAR γ activity in three PPAR CALUX cell lines upon exposure to WBM FC18.3, LA, CLA 9 *cis*-11 *trans*, and CLA 10-*trans*, 12-*cis*. PPAR α luciferase activity is expressed as a percentage of the maximum GW7647 (A) response. PPAR δ luciferase activity is expressed as a percentage of the maximum L-165.041 (B) response. PPAR γ luciferase activity is expressed as a percentage of the maximum rosiglitazone (C) response. Dotted line indicates relative induction at 10% (PC10). All concentrations of compound points showing cytotoxicity to the CALUX cells were excluded from the graphs. Data are corrected for solvent control luciferase values and are expressed as mean \pm SD in triplicate ($n = 3$).

Table 2 The lowest observed effect concentration (LOEC) of LA of purified WBM fraction FC18.3, LA, and CLA isoforms expressed as $\mu\text{g LA L}^{-1}$

	PPAR α			PPAR δ			PPAR γ		
	LOEC ($\mu\text{g LA L}^{-1}$)	LOEC relative to FC18.3	Maximum PPAR α activity of samples (%)	LOEC ($\mu\text{g LA L}^{-1}$)	LOEC relative to FC18.3	Maximum PPAR δ activity of samples (%)	LOEC ($\mu\text{g LA L}^{-1}$)	LOEC relative to FC18.3	Maximum PPAR γ activity of samples (%)
LA of FC18.3	539	1.0	75.0 \pm 3.5	3957	1.0	45 \pm 6.2	20 899	1.0	12 \pm 1.8
LA	647	1.2	69.8 \pm 3.0	6188	1.6	22.7 \pm 2.6*	11 079	0.5	19.5 \pm 3.7
CLA 9- <i>cis</i> , 11- <i>trans</i>	529	1.0	57.4 \pm 5.8*	2850	0.7	33.1 \pm 3.2*	9260	0.4	11.5 \pm 1.2
CLA 10- <i>trans</i> , 12- <i>cis</i>	5115	9.5	33.1 \pm 1.0*	13 102	3.3	24 \pm 0.7*	—	—	3.4 \pm 0.7

LOECs were expressed as microgram LA of sample per liter medium. Maximum luciferase activity of samples was expressed as a percentage of the maximum induced by PPAR reference compounds. Data are corrected for solvent control luciferase values and are expressed as mean \pm SD ($n = 3$). *: Significant difference compared with the maximum of FC18.3 PPAR activity, $P < 0.05$.

= 7.5 Hz, 2H, HOOC-CH₂), 2.07 (dt, $J = 6.9$ Hz, 4H, CH₂-CH₂-CH=), 1.66 (q, $J = 7.5$ Hz, 2H, HOOC-CH₂-CH₂), 1.43-1.26 (m, 14H), 0.91 (t, $J = 6.8$ Hz, 3H, CH₃). ¹³C{¹H} NMR (126 MHz, CDCl₃, 298 K) δ 179.23 (HOC=O), 130.24 (CH), 130.04 (CH), 128.08 (CH), 127.91 (CH), 33.89 (HOOC-CH₂), 31.54 (CH₂), 29.59 (CH₂), 29.36 (CH₂), 29.15 (CH₂), 29.08 (CH₂), 29.03

(CH₂), 27.22 (CH₂), 27.19 (CH₂), 25.64 (CH₂), 24.67 (CH₂), 22.59 (CH₂), 14.08 (CH₃) are comparable to that of a commercially obtained linoleic acid and are in agreement with the literature data of (9Z,12Z)-9,12-octadecanoic acid.⁵⁰

Linoleic acid is a major poly unsaturated fatty acid (PUFA) and an essential nutrient.⁵¹ It is of importance in a number of



structural components and physiological processes. While considered generally as beneficial, high intake may be associated with adverse effects, but no recommended daily allowance has yet been established. Being a major component of several vegetable oils, its dietary intake seems rarely insufficient.^{51,52} LA can pass through the small intestine and is packaged and transported to its targets in chylomicrons. It can also be oxidized enzymatically with its derivatives, like eicosanoids, involved in important cell signaling processes.⁵¹ We here report that LA is a major WBM-derived ligand that interacts with PPARs, which is consistent with the earlier reports of it being a PPAR ligand.^{45,46} Still, it would be difficult to envision that its levels in WBM will be responsible for the reported beneficial health effects, since its dietary intake is generally quite high.⁵¹ It is suspected that high dietary levels of LA also can lead to adverse effects.⁵² While linoleic acid is the main fatty acid in WBM, overall fungi contain low levels of fatty acids,¹⁶ which may sufficiently contribute to beneficial anti-obesogenic effects, while avoiding high energy intake counteracting its beneficial effects. In addition, these LA-induced PPAR-mediated effects may be modulated by other WBM-derived molecules. For instance, a range of polyphenols have been reported to modulate PPAR expression in a number of experimental models.⁵³ In our reporter gene assays, we tested for direct PPAR activation specifically and *in vivo* and there could be more complex cellular models modulating the effects with other WBM compounds like polyphenols, which may alter PPAR expression leading to combined effects. It can also not be totally excluded that other PPAR activating compounds may be involved. Interestingly, in particular, in the case of PPAR δ activation, a quite substantial difference was noted between the optimal fold induction by the WBM FC18.3 fraction and that by LA, being twice as high caused by the extract (Fig. 8 and Table 2), which may point towards a yet incomplete view of the PPAR activating ligands in the purified fraction, *e.g.* due to the presence of a minor component with relatively high biological activity that has a preference for PPAR δ activation.

Conclusions

A complex range of compounds is present in WBM, which could contribute to its observed anti-obesity effects.^{13,16} In this study, a stepwise fractionation was used, directed by activities observed in the bioassays to identify ligands for PPARs, which are considered important in regulating fat metabolism, and important target receptors to select ligands to combat metabolic syndrome.⁵ To our surprise, we identified a relatively common dietary fatty acid as the main ligand of PPARs in WBM, excluding other candidate molecules. Although LA is known to have relevant beneficial effects, which may be mediated through PPARs,⁵⁴ high dietary levels of LA can lead to adverse effects.⁵² While LA is the main fatty acid in WBM, overall fungi contain low levels of fatty acids,¹⁶ which may contribute to support beneficial anti-obesogenic effects, while avoiding high energy intake counteracting its beneficial

effects. However, it seems more likely that other compounds contribute to these effects. This could include a still to be discovered minor but potent compound involved in PPAR δ activation, as indicated by the relatively potent activation by the purified WBM fraction. Alternatively, receptor expression modulating compounds or compounds with completely PPAR unrelated modes of action may contribute. In further studies, we will determine if this is the only molecule responsible for the anti-obesity activities of WBM, or if there are other ones contributing, acting through different mechanisms.

Conflicts of interest

There are no conflicts to declare.

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