Conformational Plasticity and Ligand Binding of Bacterial Monoacylglycerol Lipase*

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Background: Monoacylglycerol lipases hydrolyze monoacylglycerols into free fatty acids and glycerols.
Results: Crystal structures provide the structural basis for conformational plasticity and ligand binding of a monoacylglycerol lipase.
Conclusion: Hallmark features responsible for substrate binding and selectivity are conserved across species.
Significance: The first structures of a monoacylglycerol lipase in complex with substrate analogs are presented.

Monoacylglycerol lipases (MGLs) play an important role in lipid catabolism across all kingdoms of life by catalyzing the release of free fatty acids from monoacylglycerols. The three-dimensional structures of human and a bacterial MGL were determined only recently as the first members of this lipase family. In addition to the α/β-hydrolase core, they showed unexpected structural similarities even in the cap region. Nevertheless, the structural basis for substrate binding and conformational changes of MGLs is poorly understood. Here, we present a comprehensive study of five crystal structures of MGL from Bacillus sp. H257 in its free form and in complex with different substrate analogs and the natural substrate 1-lauroylglycerol. The occurrence of different conformations reveals a high degree of conformational plasticity of the cap region. We identify a specific residue, Ile-145, that might act as a gatekeeper restricting access to the binding site. Site-directed mutagenesis of Ile-145 leads to significantly reduced hydrolysis activity. Bacterial MGLs in complex with 1-lauroylglycerol, myristoyl, palmitoyl, and stearoyl substrate analogs enable identification of the binding sites for the alkyl chain and the glycerol moiety of the natural ligand. They also provide snapshots of the hydrolytic reaction of a bacterial MGL at different stages. The alkyl chains are buried in a hydrophobic tunnel in an extended conformation. Binding of the glycerol moiety is mediated via Glu-156 and water molecules. Analysis of the structural features responsible for cap plasticity and the binding modes of the ligands suggests conservation of these features also in human MGL.

Understanding the molecular basis of enzyme substrate interactions is an important prerequisite for elucidating substrate structure-activity relationships. Structures of protein-ligand complexes are critical in delineating ligand interactions with the binding site, uncovering active site residues, and providing insight into the catalytic mechanism. In addition, they also provide valuable information on the “druggability” of the binding site, laying the basis for structure-based drug design.

Lipases catalyze the hydrolysis of ester bonds in long chain acylglycerols releasing fatty acids and glycerol. Most lipases possess a lid or cap domain that covers the active site and is able to undergo conformational changes to carry out catalysis at the lipid-water interface. The nature of these conformational changes is diverse across the lipase family and is poorly understood at an atomic level. In this study, we provide in-depth experimental insight into conformational changes and substrate binding observed in monoacylglycerol lipases (MGLs).

MGLs are a unique class of lipases (EC 3.1.1.23) because they specifically cleave monoacylglycerols (MGs) in contrast to other lipases, e.g., hormone-sensitive lipase, which accepts a broad range of substrates (triacylglycerol, diacylglycerol, cholesterol esters, and MGs), and other fungal lipases. MG-hydrolyzing lipases were first described in the 1960s, and several orthologs of MGL have been characterized over the years. The physiological function of MGL is best understood in mammals, where it has an essential role in lipid metabolism for maintaining energy homeostasis. Additionally, MGL plays an important part in mediating endocannabinoid-based signaling rendering it an important pharmacological target. In bacteria, MGLs are thought to have a role in detoxification processes because short chain MGs are highly toxic to these organisms.

Crystallographic studies of MGLs have resulted in the determination of three-dimensional structures of human MGL.

* This work was supported by the Doctoral School “DK Molecular Enzymology” Grant W901-B12 and by Project FWF P24857 funded by the Austrian Science Fund.

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VOLUME 288 • NUMBER 43

THE JOURNAL OF BIOLOGICAL CHEMISTRY

OCTOBER 25, 2013

31093

JOURNAL OF BIOLOGICAL CHEMISTRY
(hMGL) and its ortholog from a moderately thermophilic soil bacterium *Bacillus* sp. H-257 (bMGL) in free form and in complex with inhibitors (21–24). Both MGLs possess the expected α/β-hydrolase core domain harboring a catalytic triad. Additionally, the structures revealed an unexpected conservation of the overall cap architecture between hMGL and bMGL (24). Thus, bMGL can be used as an excellent model system to study the mode of action of MGLs and relate these insights to eukaryotic MGLs.

Interestingly, the cap domain of hMGL was reported to adopt open and closed conformations, whereas only an open conformation was observed for bMGL. Very little is known about the effect of these conformational changes on enzyme activity (23, 24). Similarly, the structural basis for the different substrate turnover rates shown by bMGL is also poorly understood. Reportedly, bMGL possesses higher turnover rates for MGs with shorter chain length, *i.e.* C_{10:0} and C_{12:0} compared with longer chain MGs, *i.e.* C_{18:0} and C_{18:1} (25, 26).

In this work, we describe five crystal structures of bMGL in its free form and in complex with substrate analogs and the natural substrate 1- lauroylglycerol (1-LG). We discuss their structure-function relationship, enzymatic assays, and mutations of the bacterial MGL from *Bacillus* sp. H-257. The bMGL-ligand complexes reported here represent the first experimental structures of substrate or its analogs bound to an MGL. These complex structures provide experimental evidence for conformational plasticity in the cap region of bMGL. Additionally, these structures also elucidate the molecular basis of substrate binding and help to rationalize differences in substrate turnover rates.

**EXPERIMENTAL PROCEDURES**

**Cloning, Expression, and Purification of bMGL Mutants**

Site-directed mutagenesis (I145G, I145S, and D196N) was employed to introduce single point mutations in the bMGL gene. Primers (Invitrogen) used were as follows: I145G, forward, GCCGAGGTATCTGATGCTTGCCGACGAC-TTG, and reverse, CAAGTCGACCATCCGACCATCGAT-TGCGGC; I145S, forward, GCCGAGGTATCTGATGCTTGCCGACGAC-TTG, and reverse, CAAGTCGACCATCCGACCATCGAT-TGCGGC; and D196N, forward, GCCGACGACATCGATGCTTGCCGACGAC-TTG, and reverse, GCCGAGGTATCTGATGCTTGCCGACGAC-TTG. Primers (Invitrogen) used were as follows: I145G, forward, GCCGAGGTATCTGATGCTTGCCGACGAC-TTG, and reverse, CAAGTCGACCATCCGACCATCGAT-TGCGGC; I145S, forward, GCCGAGGTATCTGATGCTTGCCGACGAC-TTG, and reverse, CAAGTCGACCATCCGACCATCGAT-TGCGGC; and D196N, forward, GCCGACGACATCGATGCTTGCCGACGAC-TTG, and reverse, GCCGAGGTATCTGATGCTTGCCGACGAC-TTG. PCRs were performed using standard protocols. After DpnI digestion, PCR products were transformed into TOP10 chemically competent *Escherichia coli* (Invitrogen). Mutations were verified by DNA sequencing, and variant enzymes were expressed and purified as described previously (24).

**Synthesis of Substrate Analogs**

The synthesis of the ligands with different alkyl chain lengths was carried out under an inert atmosphere. Two equivalents of sodium azide (1.15 g, 17.8 mmol) were added to a solution of diethyl-(3-bromopropyl)-phosphonate (2.3 g, 8.88 mmol) in 20 ml of THF/H2O (1:1), and the reaction mixture was heated for 5 h at 80 °C. After cooling to room temperature, the pH was adjusted to 10 by addition of solid NaOH. Extraction with Et2O (three times, 20 ml), drying over MgSO4, and concentrating in vacuo furnished diethyl-(3-azidopropyl)-phosphonate in quantitative yield. 500 mg (2.26 mmol) of the resulting crude product were dissolved in 2.1 ml of CH2Cl2 in a Schlenk tube. Trimethylsilyl bromide (0.92 ml, 7.0 mmol) was added, and the reaction was stirred at room temperature for 5 h. After concentration in vacuo (using a liquid nitrogen trap), crude bis(trimethylsilyl)-(3-azidopropyl)-phosphonate was produced. The yellow oil was again dissolved in 3 ml of CH2Cl2 in the same Schlenk tube. Oxalylchloride (4 eq) and a catalytic amount of N,N-dimethylformamide were added dropwise, and the reaction mixture was stirred at 40 °C for 75 min. After removal of all volatiles in vacuo (using a liquid nitrogen trap) crude dichloro-(3-azidopropyl)-phosphonate remained in the Schlenk tube. In an extra Schlenk tube, a solution of 2.26 mmol of alcohol (of different chain lengths), 0.46 ml of N,N-diisopropylethylamine, 0.41 ml of tetrazole, and 0.34 ml of diazabicyclo-[5.4.0]undec-7-ene in 2.1 ml of THF was prepared and then added in 0.5-ml portions to the other Schlenk tube containing the substrate. After overnight stirring, the resulting suspension, a solution of 4-nitrophenol (470.8 mg, 3.38 mmol) and N,N-diisopropylethylamine (0.92 ml in 2.1 ml of THF (which had been stirred overnight as well), was added. After stirring for 2 h at room temperature, the reaction mixture was concentrated using a rotary evaporator. The crude product was dissolved in CH2Cl2 (30 ml) and washed with H2O (two times, 15 ml). After back-extraction with CH2Cl2 (20 ml), the combined organic phases were washed with brine (25 ml), dried over MgSO4, concentrated, and dried in an oil pump vacuum yielding the desired alkyl-4-nitrophenyl (3-azidopropyl)phosphonate product (*e.g.* dodecyl-4-nitrophenyl (3-azidopropyl)phosphonate). The dried crude product was purified by preparative HPLC. It should be noted that a substrate analog with a 12-carbon alkyl chain actually corresponds to monomyristoylglycerol (C_{14:0}) rather than monolauroylglycerol (C_{12:0}) (Fig. 1).

**Crystallization of Substrate Analog Complexes**—For the bMGL-C_{12} ligand complex, a 0.9 mM solution of bMGL was mixed with 50 mg/ml of the C_{12} ligand dissolved in DMSO (Merck) to achieve a final concentration of 4.5 mM of the ligand. Crystallization trials were performed using the sitting drop vapor diffusion method with equal volumes of the bMGL-C_{12} complex and the Morpheus screen D4 (Molecular Dimensions, Suffolk, UK) reservoir solution containing 0.1 M MES/imidazole, pH 6.5, 12.5% w/v PEG 1000, 12.5% w/v PEG 3350, 12.5% v/v 2-methyl-2,4-pentanediol, and 0.02 M alcohols (0.2 M 1,6-hexanediol, 0.2 M 1-butanol, 0.2 M (R,S)-1,2-propanediol, 0.2 M 2-propanol, 0.2 M 1,4-butanediol, 0.2 M 1,3-propanediol). Well diffracting crystals (1.7 Å) were obtained from the drop without further optimization within 2 weeks, and no additional cryoprotectant was used for flash cooling the crystals in liquid nitrogen. For the bMGL-C_{14} ligand complex, a 0.9 mM solution of bMGL was mixed with 50 mg/ml C_{14} ligand, dissolved in 99% EtOH (Sigma) to achieve a final ligand concentration of 4.5 mM, and incubated for 1 h at 4 °C. Well diffracting crystals (1.8 Å) were obtained after 2 weeks using the hanging drop method in a crystallization condition containing 0.1 M citric acid, pH 5.2, and 18% PEG 3350. For the bMGL-C_{16} ligand complex, a 0.9 mM solution of bMGL was mixed with 50 mg/ml C_{16} ligand, dissolved in 99% EtOH (Sigma) to achieve a final ligand concent-
Crystallization of the bMGL(D196N) Variant in Complex with Substrates—An ~0.9 mM solution of bMGL(D196N) was mixed with 180 mM 1-LG (Sigma) dissolved in 99% EtOH to achieve a final protein/ligand ratio of 1:5. The protein/ligand mixture was incubated at 4 °C for 1 h. Initial crystals were obtained using the Morpheus screen, condition 4, containing 0.03 M magnesium chloride, 0.03 M calcium chloride, 0.1 M MES/imidazole, pH 6.5, 12.5% 2-methyl-2,4-pentanediol, 12.5% PEG 1000, and 12.5% PEG 3350. These crystals were used for preparing a seed stock. The optimized crystals were obtained in a drop containing 0.9 mM bMGL(D196N), 56% v/v MPD, 0.1 M HEPES, pH 6.9, and 1:1000 dilution of seed stock in a ratio of 2:2:1 respectively. To increase the probability of 1-LG being bound in the structure, we also added 1-LG as powder to these crystallization drops and soaked the crystals for 8 h. Approximately 0.9 mM bMGL(D196N) was also crystallized in the presence of 140 mM 1-(rac)-oleoylglycerol (1-OG). Crystals diffracting to 1.7 Å were obtained from the drop containing 54% v/v MPD, 0.1 M HEPES, pH 6.9, and 1:1000 dilution of seed stock in a ratio of 2:2:1, respectively.

Data Processing and Structure Refinement

Datasets were recorded at beamlines indicated in Tables 1–3. All structures were solved using molecular replacement with the structure of bMGL in its free form (PDB code 3RM3) as search template (24). Models of the ligands were created with the program MAESTRO (Maestro, version 9.3, Schrödinger, LLC, New York) and were used as input for PHENIX.elbow for creating Crystallographic Information File (CIF) library files containing bond length and angle restraints (27). The models were then subjected to rigid body and restrained refinement cycles using the program REFMAC5, followed by several iterative rounds of refinement using PHENIX (28, 29). There, water molecules were added, and the weights for the x-ray/stereochemistry and x-ray/ADP were optimized resulting in the lowest $R_{\text{free}}$ value. COOT was used to manually adjust and monitor the structures and the solvent molecules (30).

Differences in processing of datasets, $R$-values, and Ramachandran plot analysis after validation of the models using the MolProbity server are listed separately below (31). All figures displaying structures were generated using PyMOL (32).

bMGL-Ligand Complexes—The bMGL-C12 ligand complex dataset (1.7 Å) was indexed and integrated using iMosflm and scaled using Scala (33, 34). Molecular replacement was carried out using the program Phaser (35). $B$ factors of the atoms were refined anisotropically. The final model has an $R_{\text{work}}$ of 17.3% and an $R_{\text{free}}$ of 20.8%. All amino acids were in the allowed regions of the Ramachandran plot. In the final model (PDB code 4KE7), no electron density was observed for residues Thr-133–Glu-137. The bMGL-C14 ligand complex dataset (1.85 Å) was indexed and integrated using XDS and scaled using Scala (34, 36). Molecular replacement was carried out using the Balbes server (37). Model building was performed using Arp/wArp.
The three-dimensional structures of bMGL in its free form and in complex with the irreversible inhibitor PMSF also represent a snapshot of the physiological roles of MGL in hydrolyzing MG has been known for decades, yet three-dimensional structural analyses have been observed in human MGL in its free form and in the presence of different inhibitors (21–23). The three-dimensional structures of bMGL in its free form and in complex with the irreversible inhibitor PMSF also represent a snapshot of the lipase in an open conformation (24). Molecular dynamics sim-
The bMGL-C_{14} (palmitoylglycerol mimic) and bMGL-C_{16} (stearoylglycerol mimic) ligand complexes were crystallized under the same condition equally yielding crystals of space group \( P2_1 \). These structures were determined at resolutions of 1.85 and 2.2 Å, respectively. These crystals differed from the bMGL-C_{12} structure in that there were four molecules in the asymmetric unit with a different crystal packing arrangement. In all four independent molecules, electron densities for the ligands bound to active site residue Ser-97 were observed (Fig. 1, D, and E; PDB codes 4KE8 and 4KE9).

The bMGL(D196N) variant was crystallized in the presence of 1-LG (C_{12:0}) and 1-OG (C_{18:1}) yielding crystals of space group \( P2_1,2_1,2_1 \) with six molecules in the asymmetric unit. The resolutions of the datasets were 2.8 and 1.7 Å, respectively. Of both crystals, we only interpreted the electron density of the bMGL(D196N)-1-LG crystal in chain A as substrate. The electron density for the glycerol moiety of 1-LG was well defined; however, the density for carbon atoms 7 and 8 of the fatty acid chain was not observed indicating flexibility (Fig. 1F; PDB code 4KE6). Attempts to co-crystallize bMGL(D196N) with 1-OG did not result in a complex structure. Instead, a crystal form of the free enzyme in space group \( P2_1,2_1,2_1 \) was obtained containing six molecules in the asymmetric unit (PDB code 4KEA). The x-ray data collection and refinement statistics are listed in Tables 1–3.

**bMGL Shows a High Degree of Conformational Plasticity of the Cap**

The first crystal structures of hMGL in its free form and in complex with different inhibitors revealed the cap in an open conformation (21, 22). In 2011, Schalk-Hihi et al. (23) determined the structure of hMGL in complex with a reversible, noncovalent inhibitor. This structure shows significant conformational changes in the cap region compared with the lipase in its free form and to hMGL in complex with the inhibitor SAR629 (21, 22). These changes resulted in a complete closure of the binding pocket of the lipase with concomitant electrostatic differences and led to a proposal that hMGL might dissociate from the membrane during the catalytic cycle (23). bMGL crystallized in its free form and in complex with PMSF also resulted in an open conformation, herein referred to as conformation I. MD simulations of bMGL showed that the plasticity of the cap can result in the closure of both the main binding pocket and the proposed glycerol exit hole (24). So far, very little is known about the link between these conformational changes and their impact on MGL activity.

As the first remarkable feature of the different structures presented here, we highlight that bMGL captures different conformations of the cap region thus showing experimental proof for conformational changes in the cap region compared with the lipase in its free form and to hMGL in complex with the inhibitor SAR629 (21, 22). These changes resulted in a complete closure of the binding pocket of the lipase with concomitant electrostatic differences and led to a proposal that bMGL might dissociate from the membrane during the catalytic cycle (23). bMGL crystallized in its free form and in complex with PMSF also resulted in an open conformation, herein referred to as conformation I. MD simulations of bMGL showed that the plasticity of the cap can result in the closure of both the main binding pocket and the proposed glycerol exit hole (24). So far, very little is known about the link between these conformational changes and their impact on MGL activity.

**Structure Determination of Substrate and Substrate Analog Complexes**

bMGL was co-crystallized with the C_{12} ligand mimicking a C_{14} alkyl MG yielding crystals of space group \( P2_1 \), which diffracted to a resolution of 1.7 Å. Two molecules of bMGL were in the asymmetric unit, and both had the C_{12} ligand covalently bound in the active site; clear electron density for the entire ligand was observed in both chains (Fig. 1C). No electron density was observed for residues Thr-133–Glu-137 indicating their flexibility (PDB code 4KE7).

**TABLE 3**

Data collection and refinement statistics

| Contents                      | bMGL-1-LG complex | bMGL free form in \( P2_1,2_1,2_1 \) |
|-------------------------------|-------------------|--------------------------------------|
| **Data collection**           |                   |                                      |
| Beamline                      | SLS:PXIII-X06DA   |                                      |
| Wavelength                    | 1.54 Å            | 1.0 Å                                |
| Resolution                    | 73.62 to 2.8 Å    | 19.9 to 1.7 Å                       |
| Space group                   | \( P2_1,2_1,2_1 \)| \( P2_1,2_1,2_1 \)                    |
| Unit cell parameters          |                   |                                      |
| \( a, b, c \)                 | 39.19, 182.88, 244.26 Å | 39.17, 183.13, 244.70 Å |
| Total no. of reflections      | 166,183           | 159,4737                             |
| Unique reflections            | 42,482            | 195,079                              |
| \( R_{	ext{sym}} \)          | 0.096 (0.401)     | 0.051 (0.406)                       |
| Completeness                  | 94.4% (87.6%)     | 99.9% (100%)                        |
| Mean \( I/\sigma(I) \)        | 9.4 (2.6)         | 22.6 (4.4)                          |
| Multiplicity                  | 3.9 (3.1)         | 8.2 (6.6)                           |
| **Refinement statistics**     |                   |                                      |
| No. of protein atoms          | 11,010            | 115,60                               |
| No. of solvent molecules      | 39                | 934                                  |
| \( R_{	ext{work}} \)         | 20.0%             | 19.1%                                |
| \( R_{	ext{free}} \)         | 24.5%             | 21.1%                                |
| **Model geometry**            |                   |                                      |
| r.m.s.d. bonds                | 0.008 Å           | 0.006 Å                              |
| r.m.s.d. angles               | 1.360°            | 0.954°                               |
| **Ramachandran distribution**|                   |                                      |
| Most favored                  | 95.9%             | 96.2%                                |
| Additionally allowed          | 3.87%             | 3.5%                                 |
| Outliers                      | 0.22%             | 0.3%                                 |

To calculate \( R_{	ext{free}} \) 5% of the reflections were excluded from the refinement. \( R_{	ext{sym}} \) is defined as \( R_{	ext{sym}} = \Sigma_h|I(h;k) - \langle I(h;k)\rangle|/\Sigma_h|I(h;k)| \). Data in parentheses correspond to the highest resolution shell. r.m.s.d., root mean square deviation.
Ile-145 Acts as Gatekeeper for the Substrate Entrance Tunnel and the Glycerol Exit Hole

The different conformations observed provide an experimental corroboration for conformational plasticity of the bMGL cap region (Fig. 2). Therefore, we wanted to study the residues involved in mediating these conformational changes. A comparison between the super-open conformation (as observed in the free form of bMGL(D196N) chain A), conformation I (as observed in free bMGL, bMGL-PMSF complex, and the bMGL-C12 complex, and four molecules of free bMGL(D196N)), conformation II (as observed in two molecules of bMGL-C14 and bMGL-C16 complex structures and one molecule of free bMGL(D196N)), and conformation III (as observed in chains C and D of the bMGL-C14 and bMGL-C16 complex structures) shows that the position and conformation of Ile-145 differ extensively (Fig. 2, D and E). Inspection of Ile-145 in conformations I and II shows that the backbone Ca atom position of Ile-145 differs by about 2.7 Å accompanied by 1.8- and 5.1-Å movements of the CG2 and the CD1 side chain methyl groups, respectively. The geometric centers of Ile-145 differ by 3.0 Å (Fig. 3A). This results in a partly restricted conformation of the binding pocket in conformation II. Again, in conformation III the residue playing a crucial role in the observed changes is Ile-145. The CG2 and CD1 methyl groups are flipped 4.1 and 9.6 Å away from the open conformation, respectively. The geometric centers of Ile-145 differ by 4.9 Å (Fig. 3B). In the super-open conformation, the side chain of Ile-145 points into the solvent region without engaging in any crystal contacts (Fig. 3, C and D). The geometric center of Ile-145 in the super-open and conformation III differs by 7.6 Å (Fig. 3C). The closure of the exit hole in conformation III is mediated by a 6.4-Å movement of the main chain Ca of Ile-145 compared with the super-open conformation (Fig. 3D). In the bMGL-C14 complex, the movement of Ile-145 also restricts the orientation of the azide headgroup of the C14 ligand bound to Ser-97. In conformation III, Ile-145 engages in close contacts with the ligand, which results in positioning of the azide headgroup deep within the binding pocket to avoid steric clashes with Ile-145.

Thus, our crystal structures clearly show that Ile-145 plays a crucial role in engaging in conformational changes, modifying access to the substrate-binding site of bMGL, and determining open and closed states of the exit hole (Fig. 3). These observations pave the way for an intriguing set of questions relating to critical residues and the functional relevance of these different
We therefore investigated whether this residue also influences the catalytic activity of the lipase. Two new variants of bMGL were generated by replacing Ile-145 with Ser and Gly, respectively. The variants were tested for MG hydrolase activity using 1-lauroyl-rac-glycerol (C12:0) and 1-oleoyl-rac-glycerol (C18:1). As reported before, WT-bMGL had a more than 2-fold higher activity toward the medium chain 1-LG compared with the longer chain 1-OG (25, 26). Compared with the wild-type protein, decreased activity levels were observed for the bMGL I145G variant against 1-OG and 1-LG (82% remaining activity for both substrates). Interestingly, the I145S variant harboring a polar side chain showed an even more drastic loss of activity toward these substrates (38 and 62% for 1-OG and 1-LG, respectively) (Fig. 3E). This shows that increasing the flexibility of this loop region and concomitantly removing the hydrophobic side chain reduces activity. The introduction of a polar side chain has an even bigger effect indicating that a hydrophobic side chain is required at this position for optimal activity. One might speculate that such a hydrophobic residue aids in recruitment of the substrate, which itself harbors a long hydrophobic alkyl chain.

Next, we looked at the size of the substrate binding pocket in more detail. Analysis of the binding cavity revealed that the movement of Ile-145 from conformation I to conformation III squeezes the space within the binding pocket (39). Thus, it is tempting to speculate that conformation III (Fig. 3F) could limit the cavity to bind only MG and not diacylglycerol or triacylglycerol as bMGL shows no activity toward these substrates (26).

Identification of the Binding Mode of the Substrate and Importance of Glu-156 for Glycerol Binding

The bMGL-C12, bMGL-C14, and bMGL-C16 complex structures show that the hydrophobic nature of the substrate binding pocket plays a major role in stabilizing the fatty acid moiety of the substrate. Representative of these contacts, the side chains of residues from both the α/β-hydrolase core and the cap region, including Phe-29, Ile-125, Ile-128, Leu-142, Leu-
167, Leu-170, Met-174, and Val-198, form hydrophobic contacts with the alkyl chain of the C_{12}-ligand bound at the active site in the bMGL-C_{12} complex (Fig. 4A). The otherwise flexible fatty acid chain of an MG substrate is observed in a single conformation in the crystal structure. The backbone NH groups of Phe-29 and Met-98 form hydrogen bonds with the O2 oxygen atom of the phosphonate group of the C_{12} ligand, highlighting that these residues form the oxyanion hole during the course of MG hydrolysis (Figs. 4A and 5C). In the free form of the lipase, a water molecule occupies this position (24).

bMGL has been reported to have a preference for C_8-C_{14} chains with 1-LG (C_{12}) identified as the best substrate among those tested. Our results are in agreement with these previous reports (Fig. 3E) (25, 26). The structures reported here reveal the impact of chain length on the interactions of substrate with the binding site. The bMGL-C_{12} ligand complex shows the aliphatic chain of the ligand in an almost linear conformation protruding toward the surface (Figs. 4B and 1C). In the bMGL-C_{14} complex, a bend in the aliphatic chain allows the longer chain to be accommodated in the binding pocket (Figs. 4C and 1D). The terminal carbon atoms of both the C_{14} and C_{16} alkyl chains are located at approximately the same distance from the surface of the protein, close to residues that line the entrance of the binding pocket (Fig. 4C). The C_{16} ligand is also fully accommodated in the binding pocket, despite the four additional CH_2 groups, due to alkyl chain bending (Figs. 4D and 1E). The necessity for bending might explain the lower substrate turnover rate of bMGL with respect to MGS with a longer fatty acid chain.

Next, we analyzed the binding mode of the glycerol moiety of the MG substrate. To stall the catalytic activity, yet retain a WT-like architecture of the catalytic center, we generated an inactive variant of bMGL by introducing an D196N point mutation. The structure of the bMGL(D196N)-1-LG complex shows the interactions between the glycerol moiety of an MG substrate and residues at the bottom of the substrate binding pocket. The carbonyl carbon of 1-LG is located 3.1 Å away from the oxygen OG of the nucleophilic Ser-97 (Fig. 5A). The glycerol moiety of 1-LG forms contacts with the side chain of Glu-156 and a water molecule. The hydroxyl group of Ser-35 and the backbone nitrogen of Gly-28 are also engaged in interactions with the glycerol moiety via this water molecule (Fig. 5B).

Interestingly, two water molecules are positionally conserved in the other bMGL structures (Fig. 5D). The fact that only one water was observed in the bMGL(D196N)-1-LG complex structure might be due to the lower resolution of that dataset.

bMGL Complexes Provide Detailed Insights into the First Steps of the Lipolytic Reaction

The catalytic mechanism of bMGL is expected to follow the typical hydrolysis mechanism of α/β-hydrolases involving the catalytic triad residues Ser-97, Asp-196, and His-226 and the oxyanion hole forming residues Met-98 and Phe-29 (Fig. 5A) (24, 40). Asp-196 forms a hydrogen bond to a histidine imidazole nitrogen thus enabling His-226 to act as the general base and to accept the proton from the catalytic serine. The deprotonated serine acts as a powerful nucleophile attacking the carbonyl carbon of the MG substrate. The spatial arrangement of the catalytic triad residues in the free enzyme structure is consistent with this general reaction scheme (24). In the D196N mutant, in addition to a loss of charge, the tautomeric state of the histidine side chain is likely to be altered. The reaction is stalled before the nucleophilic attack with the entire ligand in the active site representing the Michaelis complex. The distance between Ser-OG and the carbonyl carbon of the 1-LG substrate is 3.1 Å (Fig. 5A). In analogy to serine proteases, the main chain atoms of Met-98 and Phe-29 form the oxyanion hole, which would stabilize the tetrahedral intermediate formed in the course of MG hydrolysis. Accordingly, the main chain NH groups are in hydrogen bonding distance from the 1-LG carbonyl-oxygen in the structure (Fig. 5A).

We synthesized p-nitrophenol esters of alkyl phosphonic acids and used them as substrate analogs with different alkyl chain lengths. These compounds react irreversibly with the nucleophilic serine of α/β-hydrolases and are used frequently to capture the tetrahedral reaction intermediates of lipases (41–45). Consequently, the inhibitor complexes go one step further along the reaction trajectory compared with the D196N 1-LG
complex; Ser-97 exerted its nucleophilic attack on the phosphorus-atom of the MG-mimicking ligand, and then the reaction is stalled. The p-nitrophenol moiety is released and no longer present in the active site (Fig. 5C). The irreversibly and covalently bound ligand is captured in the bMGL/H18528C12, bMGL/H18528C14, and bMGL/H18528C16 complex structures (distance between Ser-OG and the phosphorus atom, 1.6 Å) mimicking the tetrahedral intermediate of carboxylic acid ester hydrolysis. The ligands also form hydrogen bonds to the oxyanion hole residues Met-98 and Phe-29. The distances between backbone hydrogen donor nitrogens and phosphonate oxygens are 2.8 and 2.9 Å (Fig. 5C). The next steps in the catalytic reaction of an MGL substrate would be protonation and release of the glycerol moiety. His-226 is very likely to act as the proton donor. Following glycerol release, the ester carbonyl would likely be held in place by Met-98 and Phe-29 in the acyl-enzyme intermediate. In the free state of bMGL, a further water molecule is located in hydrogen bonding distance from His-226 NE2 that could occupy a similar position to a water molecule required for fatty acid release. The position and orientation of the ligands in our structures are consistent with such a prototypical hydrolysis mechanism.

Conservation of Structural Features of bMGL in Human MGL

The structure of bMGL in its free form revealed structural conservation of the α/β-hydrolase core region and unexpected evolutionary conservation of the cap architecture between human and bacterial proteins (24). The structures of bMGL in complex with substrate and the analogs reported here shed light on the conservation of additional structural features important for the hydrolytic action of these two proteins.

Binding Pocket Hydrophobicity—The lipophilic potential of bMGL calculated using VASCo indicated that the hydrophobicity of the protein is largely restricted to the main substrate binding pocket (24, 39). The structures of bMGL in complex with 1-LG, C12, C14, and C16 ligands corroborate the importance of the hydrophobicity of the binding pocket in the otherwise polar and water-soluble protein (Fig. 4). The hydrophobic interactions between the side chains of the residues lining the
pocket and the aliphatic carbon chain of the ligand provide an ideal environment for the binding of the fatty acid moiety of an MG. The structures of hMGL and bMGL show that the hydrophobicity of the binding pocket is conserved between the two lipases (21–23). It suggests that despite the low sequence identity of only 17%, the conservation of the hydrophobicity of the substrate binding pocket is one of the essential factors for MGL function.

Glycerol Binding—Although structures of hMGL have been determined in free form and in complex with inhibitors, very little is known about the glycerol binding pocket of MGLs. The structure of bMGL in complex with 1-LG provides first insights into the important role of Glu-156 and a water molecule for binding the glycerol moiety of the substrate. Our bMGL structures determined in free form and in complex with ligand show that one water molecule is directly involved in binding the glycerol moiety, but two are positionally conserved in all the other bMGL structures (Fig. 5, B and D). In the human ortholog, docking studies indicated that Tyr-194 interacts with the glycerol moiety. This represents a difference to bMGL where we identified Glu-156 as a key residue for glycerol binding (22). hMGL also harbors conserved water molecules in the glycerol binding pocket. In the structure of hMGL in its free form (PDB code 3HJU (21)), Tyr-194 forms a hydrogen bond with a water molecule located at a distance of 2.7 Å (Fig. 5E) (22). In another structure (PDB code 3W8), a water molecule in a similar position forms a hydrogen bond with this tyrosine residue at a distance of 2.5 Å. In the hMGL structure in the closed conformation, two water molecules engage in hydrogen bonding interactions with Tyr-194 (PDB code 3PE6 (23)). Interestingly, these water molecules are located at different positions compared with the other two hMGL structures. This could be due to the fact that the ligand co-crystallized with the protein extends into the glycerol binding pocket. Hence, water molecules probably play a role in glycerol binding in human as well as bacterial MGLs.

Isoleucine Residues Undergo Conformational Changes in the Cap—bMGL structures reveal that Ile-145 is the major residue undergoing conformational changes in the cap region acting as a gatekeeper. These movements affect the access to the substrate binding pocket and the closure of the exit hole. The structure of hMGL determined in the closed conformation with a noncovalent inhibitor possesses strikingly similar properties. The closure of the exit hole in this complex coincides with movements of Ile-179 (23). This residue is located at an equivalent position to Ile-145 in bMGL with the backbone atoms lining the glycerol exit hole (Fig. 6A).Analysis of the three-dimensional structure of hMGL reveals that both the side chain and the main chain of Ile-179 change conformation to open and close the proposed glycerol exit hole (Fig. 6B).

In summary, the structures of bMGL in complex with substrate and substrate analogs reveal the mode of substrate binding in MGLs and capture different stages of the lipolytic reaction. In complex structures with substrate analogs of different chain lengths, bending of longer alkyl chains is observed, which might contribute to faster turnover rates of MG substrates with short chains. The bMGL structures shed light on the stochastic equilibrium between open and restricted cap conformations. Analysis of the binding pocket leads to the hypothesis that restricted conformations result in substrate selectivity toward MG only. It remains to be seen whether these cap conformations are essential for catalysis and release of reac-
tion products. These questions might be addressed in future studies by using a combined approach of time-dependent soak-freeze x-ray crystallography, site-directed spin labeling, and NMR spectroscopy.

Acknowledgments—We thank Dr. Georg Steinkellner for providing the program Casox for cavity calculations. We are also grateful for the support in data collection at the Synchrotrons ESRF (Grenoble, France) and SLS (Villigen, Switzerland).

REFERENCES

1. Thornton, J. M., Todd, A. E., Milburn, D., Borkakoti, N., and Orengo, C. A. (2000) From structure to function: approaches and limitations. Nat. Struct. Biol. 7, 991–994

2. Whittle, P. J., and Blundell, T. L. (1994) Protein structure-based drug design. Annu. Rev. Biophys. Biomol. Struct. 23, 345–375

3. Brzozowski, A. M., Derewenda, U., Derewenda, Z. S., Dodson, G. G., Law- son, D. M., Turkenburg, J. P., Bjorkling, F., Hug-Jensen, B., Patkar, S. A., and Thim, L. (1991) A model for interfacial activation in lipases from the structure of a fungal lipase-inhibitor complex. Nature 351, 491–494

4. Derewenda, U., Brzozowski, A. M., Lawson, D. M., and Derewenda, Z. S. (1992) Catalysis at the interface, the anatomy of a conformational change in a triglyceride lipase. Biochemistry 31, 1532–1541

5. Liu, L., Gao, C., Lan, D., Yang, B., and Wang, Y. (2012) Molecular basis for substrate selectivity of a mono- and diacylglycerol lipase from Malassezia globosa. Biochem. Biophys. Res. Commun. 424, 285–289

6. Xu, T., Liu, L., Hou, S., Xu, J., Yang, B., Wang, Y., and Liu, J. (2012) Crystal structure of a mono- and diacylglycerol lipase from Malassezia globosa reveals a novel lid conformation and insights into the substrate specificity. J. Struct. Biol. 178, 363–369

7. Fredriksson, G., Strålfors, P., Nilsson, N. O., and Belfrage, P. (1981) Hormone-sensitive lipase of rat adipose tissue. Purification and some properties. J. Biol. Chem. 256, 6131–6130

8. Pope, J. L., Askins, R. E., and McPherson, J. C. (1962) The specificity of an intestinal lipase for monoglycerides. Fed. Proc. 21, 259

9. Côté, K., Dhouib, R., Douchet, I., Chahinian, H., de-Caro, A., Carrière, F., and Canaan, S. (2007) Characterization of an exported monoglycerol lipase from Mycobacterium tuberculosis that plays a role in macrophage function. J. Biol. Chem. 282, 408, 417–427

10. Dhouib, R., Laval, F., Carrière, F., Daffé, M., and Canaan, S. (2010) A monoglycerollipase from Mycobacterium smegmatis involved in bacterial cell interaction. J. Bacteriol. 192, 4776–4785

11. Sakiyama, T., Yoshimi, T., Miyake, A., Umeoka, M., Tanaka, A., Ozaki, S., and Nakanishi, K. (2001) Purification and characterization of a monoglycerol lipase from Pseudomonas sp. LP7315. J. Biosci. Bioeng. 91, 27–32

12. Karlsson, M., Tornqvist, H., and Holm, C. (2000) Expression, purification, and characterization of histidine-tagged mouse monoglyceride lipase from baculovirus-infected insect cells. Protein Expr. Purif. 18, 286–292

13. Heier, C., Taschler, U., Radner, F., Oberer, M., Wolinski, H., Natter, K., Kohlwein, S. D., Leber, R., and Zimmermann, R. (2010) Identification of YjciE as functional ortholog of mammalian monoglyceride lipase in the yeast Saccharomyces cerevisiae. Biochem. Biophys. Acta 1801, 1063–1071

14. Taschler, U., Radner, F., Heier, C., Schreiber, R., Schweiger, M., Schoiswohl, G., Preis-Landl, K., Jaeger, D., Reiter, B., Koefeler, H. C., Wajciechowski, J., Theussl, C., Penninger, J. M., Lass, A., Haemmerle, G., Zechner, R., and Zimmermann, R. (2011) Monoglyceride lipase deficiency in mice impairs lipolysis and attenuates diet-induced insulin resistance. J. Biol. Chem. 286, 17467–17477

15. Dinh, T. P., Kathuria, S., and Piomelli, D. (2004) RNA interference suggests a primary role for monoglycerol lipase in the degradation of the endocannabinoid 2-arachidonoylglycerol. Mol. Pharmacol. 66, 1260–1264

16. Labar, G., Wouters, J., and Lambert, D. M. (2010) A review on the monoglycerol lipase: at the interface between fat and endocannabinoid sig-
Structures of Bacterial Monoglyceride Lipase

36. Kabsch, W. (2010) Xds. Acta Crystallogr. D Biol. Crystallogr. 66, 125–132
37. Long, F., Vagin, A. A., Young, P., and Murshudov, G. N. (2008) BALBES: a molecular replacement pipeline. Acta Crystallogr. D Biol. Crystallogr. 64, 125–132
38. Perrakis, A., Morris, R., and Lamzin, V. S. (1999) Automated protein model building combined with iterative structure refinement. Nat. Struct. Biol. 6, 458–463
39. Steinkellner, G., Rader, R., Thallinger, G. G., Kratky, C., and Gruber, K. (2009) VASCo: computation and visualization of annotated protein surface contacts. BMC Bioinformatics 10, 32
40. Schrag, J. D., and Cygler, M. (1997) Lipases and α/β-hydrolase fold. Methods Enzymol. 284, 85–107
41. Colton, I. J., Yin, D., Grochulski, P., and Kazlauskas, R. J. (2011) Molecular basis of chiral acid recognition by Candida rugosa lipase: x-ray structure of transition state analog and modeling of the hydrolysis of methyl 2-methoxy-2-phenylacetate. Advanced Synthesis and Catalysis 353, 2529–2544
42. Egloff, M. P., Marguet, F., Buono, G., Verger, R., Cambillau, C., and van Tilbeurgh, H. (1995) The 2.46 Å resolution structure of the pancreatic lipase-colipase complex inhibited by a C11 alkyl phosphonate. Biochemistry 34, 2751–2762
43. Mannesse, M. L., Boots, J. W., Dijkman, R., Slotboom, A. I., van der Hieden, H. T., Egmond, M. R., Verheij, H. M., and de Haas, G. H. (1995) Phosphonate analogs of triacylglycerols are potent inhibitors of lipase. Biochim. Biophys. Acta 1259, 56–64
44. Marguet, F., Cudrey, C., Verger, R., and Buono, G. (1994) Digestive lipases: Inactivation by phosphonates. Biochim. Biophys. Acta 1210, 157–166
45. Susanni-Etzerodt, H., Schmidinger, H., Riesenhuber, G., Birner-Gruenberger, R., and Hermetter, A. (2006) A versatile library of activity-based probes for fluorescence detection and/or affinity isolation of lipolytic enzymes. Chem. Phys. Lipids 144, 60–68
46. Baker, N. A., Sept, D., Joseph, S., Holst, M. J., and McCammon, J. A. (2001) Electrostatics of nanosystems: Application to microtubules and the ribosome. Proc. Natl. Acad. Sci. U.S.A. 98, 10037–10041