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Lipase catalyzed oxidations in a sugar-derived Natural Deep Eutectic Solvent

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58 Abstract

59 Chemoenzymatic oxidations involving the CAL-B/H₂O₂ system was developed in a sugar derived 60 Natural Deep Eutectic Solvent (NaDES) composed by a mixture of glucose, fructose and sucrose. Good to excellent conversions of substrates like cyclooctene, limonene, oleic acid and stilbene to 61 cyclohexanone to its corresponding 62 their corresponding epoxides, lactone and 2-63 phenylacetophenone to its corresponding ester, demonstrate the viability of the sugar NaDES as a 64 reaction medium for epoxidation and Baever-Villiger oxidation.

66 Keywords

67 Chemoenzymatic oxidations, green solvents, Natural Deep Eutectic Solvents, Epoxidation, Baeyer 68 Villiger, Lipase

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71 **1. Introduction**

72 Oxidation reactions have always been a major area of research due to their tremendous industrial 73 applications. However, several oxidation processes present sustainability issues from the point of 74 view of oxidants, catalysts and solvents used. (Cavani & Teles 2009) For example, a peracid is 75 often used as the oxidizing agent, (Swern 1949) but transportation and storage of organic peracids leads to significant safety issues and costs; when achievable, molecular oxygen or air are for sure 76 77 the ideal oxidants, with hydrogen peroxide as the second-best choice, in terms of atom economy 78 and applicability to various oxidation systems. (O. Burek et al. 2019) In this context, enzymes, that 79 can work in sustainable solvents with mild oxidants, can contribute to increase the greenness of the 80 oxidation reactions. (Niku-Paavola & Viikari 2000; Constable et al. 2007; Gorke et al. 2008; Kotlewska et al. 2011; Silva et al. 2011; Hollmann et al. 2011; Drożdż et al. 2015; Qin et al. 2015; 81 82 Yin et al. 2015; Yang & Duan 2016; García et al. 2018)

In fact, a very interesting system for obtaining peracids in situ is the chemoenzymatic system 83 84 lipase/H₂O₂ that continuously forms the peracids through a lipase-catalyzed perhydrolysis of 85 carboxylic acids or their esters. (Björkling et al. 1990; Yadav & Devi 2002; Busto et al. 2010) A 86 broad range of hydrolases has been investigated for the peracid formation and among them the 87 lipase B from Candida Antarctica (CAL-B), immobilized onto an acrylic resin (Novozyme 435) is 88 the most reactive. (Ortiz et al. 2019) This system has been successfully applied to both epoxidations 89 of alkenes (Prileshajev-epoxidation) and Baeyer-Villiger (B-V) oxidations (Scheme 1). (Lemoult et 90 al. 1995; Aouf et al. 2014)

91 Epoxides are fundamental intermediates in organic synthesis but, despite their relevance, their 92 industrial synthesis is scarcely sustainable (both environmentally and economically). Epoxidation of some natural products is industrially carried out by the Prileshajev-epoxidation (epoxidation of an 93 94 alkene with a peracid) using either preformed or in-situ-generated short chain peroxy acids. (Rüsch 95 gen. Klaas & Warwel 1999; Hilker et al. 2001) Nevertheless, the need for a strong acid to catalyze 96 peroxy acid formation in this process can result in unsatisfactory selectivity and undesiderable side 97 reactions via oxirane ring opening, leading to diols, hydroxyesters, and dimers. Prileshajev-98 epoxidation can be chemoenzymatically carried out on various substrates with CAL-B, a carboxylic 99 acid as precursor of the peracid, and H₂O₂ as oxidant (Scheme 1); this method allows an improvement in terms of sustainability, mild reaction conditions, limited side products and use of 100 101 less toxic reagents. (Niku-Paavola & Viikari 2000; Moreira & Nascimento 2007; Silva et al. 2011; 102 Hollmann et al. 2011)

The B-V oxidation is a very well-known and useful reaction for the synthesis of esters and lactones starting from ketones, important building blocks in pharmaceutical and polymer synthesis. (Renz & Meunier 1999; Brink et al. 2004; Woodruff & Hutmacher 2010) Peracids such as metachloroperbenzoic acid or peracetic acid are used as stoichiometric reagents, but also various catalytic methods that use metals have been studied. (Strukul 1998; Ma et al. 2014) Protocols based on B-V monooxygenases have also been developed, but given their need for oxygen, cofactor NADPH and their intrinsic low stability, they are considered unpractical. (Alphand et al. 2003;
Leisch et al. 2011; Balke et al. 2012) Nevertheless, the simple chemoenzymatic method based on
CAL-B, used for epoxidation of alkenes described above, has been also applied to this kind of
oxidations (Scheme 1). (Lemoult et al. 1995; Ríos et al. 2007; Rios et al. 2008)

Green solvents exploited in chemoenzymatic oxidation reactions can be categorized into two main 113 114 groups: i) water and ii) non-aqueous solvents like ionic liquids, (Moniruzzaman et al. 2010; 115 Elgharbawy et al. 2020) supercritical fluids and fluorinated solvents. (Hobbs & Thomas 2007) Despite their interesting properties and application possibility, ionic liquids suffer from several 116 drawbacks like cost, toxicity, low biodegradability, complexity in preparation and handling. 117 118 (Samorì et al. 2015; Lei et al. 2017; B. Wang et al. 2017) Deep Eutectic Solvents (DESs), described for the first time by Abbott et al., (Abbott et al. 2001) are low melting mixtures based on a 119 combination of readily-available and inexpensive components, like quaternary ammonium salts as 120 hydrogen bond acceptors (HBA), and acids, amides, amines, carbohydrates and alcohols as 121 122 hydrogen bond donors (HDB). They are liquid at or below 100 °C, thanks to H-bond interactions 123 between the single components that create specific supramolecular structures. The number and the spatial positions of hydrogen atoms in the donor and acceptor groups, available for hydrogen 124 bonding, influence the formation and stability of the DES itself. (Nkuku & LeSuer 2007; Zhang et 125 al. 2012; Paiva et al. 2014; Smith et al. 2014; Tommasi et al. 2017; Samorì et al. 2019) Dai et al. 126 reported numerous preparations of Natural Deep Eutectic Solvents (NaDESs) by using plant 127 metabolites. Interestingly, when water is added to the mixtures, in different proportions according to 128 the NaDES, it can be incorporated into this structure and becomes strongly bound, reducing the 129 130 viscosity of the NaDES while retaining its original characteristics. (Dai et al. 2013)

The chemoenzymatic oxidation systems described above (Scheme 1) have been studied in several 131 solvents, including DESs (T. Gorke et al. 2008; Kotlewska et al. 2011; Durand et al. 2012; Drożdż 132 133 et al. 2015; Yang & Duan 2016; Zhou et al. 2017; Ranganathan et al. 2017; Gotor-Fernández & Paul 2019; Ma et al. 2019), in which it has been demonstrated that the enzymatic activity, stability, 134 135 and selectivity can be enhanced. (Zhou et al. 2017; Ülger & Takaç 2017; Oh et al. 2019; Guajardo et al. 2019; Gotor-Fernández & Paul 2019; Nian et al. 2020). Among the various NaDES proposed 136 137 by Dai et al. we focused on the only one sugar-derived and chlorine-free combination, composed by glucose, fructose sucrose and water (1:1:1:11), that we called with the acronym GFS. To the best of 138 139 our knowledge the lipase/H₂O₂ system was never reported in a solvent like GFS and, following our interest in biocatalysis in sustainable reaction media, (Galletti et al. 2007) herein we report on its 140 application in the epoxidation of alkenes and B-V oxidation of ketones. 141

142

143 **2. Materials and methods**

144 <u>2.1 Material</u>: all chemicals and solvents were purchased from Sigma-Aldrich or Alfa Aesar and
 145 used without any further purification.

- 146 CAL-B (Lipase B from *Candida antarctica*) immobilized on Immobead 150, recombinant from
 147 yeast, 4000 U/g was used.
- <u>2.2 DESs preparation</u>: the components were mixed with the appropriate stoichiometric ratios,
 heated at about 80-90 °C (120 °C for GFS) and magnetically stirred until homogeneous liquid was
- obtained; for GFS, distilled water (up to 30 wt %) was then added to get a homogeneous colorless liquid phase. All the DESs were cooled to rt (20 °C) before the use and stored in the fridge (4 °C).
- 152 2.3 Representative procedure for enzymatic epoxidations of alkenes: in a 4-mL vial, the
- immobilized CAL-B (amounts reported in Tables 1-4 and Scheme 4) and 400 mg of DES (200 mg
- 154 for 1d, 800 mg for 1e) were weighted, followed by the addition of 1.6 mmol of alkene (0.8 mmol
- 155 for **1f**), carboxylic acid (amounts reported in Tables 1-4 and Scheme 4) and 1 equivalent (eq) of
- 156 H_2O_2 (30% aqueous solution). For entries 17 and 18 in Table 1, H_2O_2 has been added in 4 aliquots
- 157 in 4 h, for substrate **1f** 1.5 eq has been used.
- 158 The vial was heated at 45 °C (or rt, 20 °C, for **1d**) for various reaction times, then crudes were 159 extracted with cyclohexane or ethyl acetate and analyzed by GC-MS. Extraction residues were

160 checked after derivatization by silylation for the presence of other by-products (see section 2.5). 161 Conversions were calculated as ratios between products areas and total areas. ¹H and ¹³C NMR 162 spectra of some products have been acquired after purification of the crude by flash-column 163 chromatography (see section 2.6), some isolated yields are also reported in the Tables. All products 164 are known, they were recognized by comparison with standards or through mass spectra matching 165 to what reported in NIST database. Formation of byproducts was checked by GC-MS and NMR.

- 166 <u>2.4 Representative procedure for B-V oxidation of ketones</u>: in a 4-mL vial, the immobilized CAL-B
- 167 (amounts in Tables 3-4) and 400 mg of GFS for **3a**, **3b**, **3c**, (700 mg of GFS for **3d**) were weighted, 168 followed by the addition of 0.8 mmol of ketone (0.4 mmol for **3d**), carboxylic acid (amounts in
- Tables 3-4) and 1 eq of H_2O_2 (30% aqueous solution), different amounts of H_2O_2 are reported in Table 3. The vial was heated at 45 °C or at kept at rt, 20 °C, for various reaction times, then crudes were extracted with ethyl acetate and analyzed by GC-MS as described above. ¹H and ¹³C NMR spectra of some purified products have been acquired after purification (see section 2.6) of the crude
- 173 by flash-column chromatography, some isolated yields are also reported in Tables 3-4.
- 174 <u>2.5 Silylation procedure</u>: 50 μ L of silylating agent *N*,*O*-bis(trimethylsilyl)trifluoroacetamide and 175 1% chlorotrimethylsilane, (BSTFA + 1% TMCS), 100 μ L of CH₃CN and 20 μ L of pyridine were 176 added to 1–10 mg of sample into a GC-MS vial. The vial was heated at 60–80 °C for 30-40 min. 177 The sample was then diluted with CH₃CN before the injection.
- <u>2.6 Purification procedure of selected products</u>: reaction mixtures were extracted with ethyl acetate
 or cyclohexane then washed with a NaHCO₃ solution to remove the octanoic acid (OA). After
 evaporating the solvent, the crude was purified by flash chromatography. The fractions containing
 the product were mixed, the solvent evaporated, and the purified products were analyzed by GC-MS
 and NMR (See spectra in supplementary information, SI).
- 2.7 Instrumentation: GC-MS analysis of epoxides and ester 4d were performed using an Agilent HP 183 6850 gas chromatograph connected to an Agilent HP 5975 quadrupole mass spectrometer. Analytes 184 were separated on a HP-5MS fused-silica capillary column (stationary phase 5%-Phenyl)-185 186 methylpolysiloxane, 30 m, 0.25 mm i.d., 0.25 µm film thickness), with helium as the carrier gas (at constant pressure, 36 cm s⁻¹ linear velocity at 200 °C). Mass spectra were recorded under electron 187 ionization (70 eV) at a frequency of 1 scan s⁻¹ within the 12–600 m/z range. The injection port 188 temperature was 250 °C. The temperature of the column was kept at 50 °C for 5 min, then increased 189 from 50 to 250 °C at 10 °C min⁻¹ and the final temperature of 250 °C was kept for 12 min. 190
- 191 GC-MS analysis of Baeyer-Villiger products (except **4d**) were performed using an Agilent 7820A 192 gas chromatograph connected to an Agilent 5977E quadrupole mass spectrometer. Analytes were 193 separated on a DB-FFAP polar column (30 m length, 0.25 mm i.d., 0.25 μ m film thickness), with 194 helium flow of 1 mL min⁻¹. Mass spectra were recorded under electron ionization (70 eV) at a 195 frequency of 1 scan s⁻¹ within the 29–450 m/z range. The injection port temperature was 250 °C.
- 196 The temperature of the column was kept at 50 °C for 5 min, then increased from 50 to 250 °C at 10 197 °C min⁻¹ and the final temperature of 250 °C was kept for 15 min.
- ¹H NMR spectra were recorded on Varian 400 (400 MHz) spectrometers. ¹³C NMR spectra were
 recorded on a Varian 400 (100 MHz) spectrometers. Chemical shifts were reported in ppm from
 trimethylsilane (TMS) with the solvent resonance as the internal standard (deuterochloroform: 7.26
 ppm).
- 202

203 **3. Results and discussion**

204 <u>3.1 Alkenes epoxidation</u>

We studied the chemoenzymatic epoxidation of various alkenes (focusing for each of them on the enzyme amount, peracid precursors and H_2O_2 amount and additions, and reaction time): cyclic alkenes (Table 1), poorly-reactive stilbene (Table 2) and oleic acid (Scheme 2).

- 208
- 209 *3.1.1 Cyclic alkenes* **1** *a*-*d*

210 Studying CAL-B mediated chemoenzymatic epoxidation in various NaDESs, Zhou et al. showed that amine-based DESs (i.e. choline chloride-urea, 1:2 molar ratio, called Reline) significantly 211 212 reduced the stability of CAL-B in a wide temperature range whereas the polyol-based ones increased it. (Zhou et al. 2017) For this reason, we focused our initial experiments on polyol-based 213 NaDESs. Cyclohexene 1a (Table 1, entry 1) was epoxidized with immobilized CAL-B (100 mg per 214 215 1.6 mmol of alkene), octanoic acid OA (one eq respect to the alkene), and H₂O₂ (one eq respect to the alkene) in choline chloride-sorbitol (ChCl-Sorb), 1:1 molar ratio (400 mg) (similarly to Zhou et 216 al). We observed a complete conversion of the starting material but a very low selectivity towards 217 the epoxide; in fact the most of epoxide was converted into the chlorinated by-product and the diol 218 219 after 20h. This unexpected result prompted us to turn our attention towards chloride-free, sugarbased NaDESs as the GFS. We tested both solvents (ChCl-Sorb and GFS) in the same conditions 220 221 with more easily detectable cyclodocedene **1b** and we observed that GFS gave better conversions (Table 1, entries 2 and 3); the same held true for other cyclic substrates (cyclooctene 1c and 222 223 limonene 1d) (see in SI Table S1, entries 1 and 5, and Table S2, entry 1). So, we decided to test 224 various substrates to check the viability of the system.

When Z/E mixtures were used in the starting alkene (as in **1b**), no diastereoselectivity was observed and the final product diastereomeric ratio reflected the diastereomeric distribution in the reagent.

OA resulted the most reactive acid precursor under our conditions (Table 1, entries 9-12), 227 228 confirming the literature results, and its amount can be significantly lowered from 1 eq to 0.1 eq with all the substrates (Table 1, entries 6, 8, 13, 14, 18). Considering aliphatic acids with different 229 chain lengths, butanoic acid BA (Table 1, entry 10) gave very good results on 1c while acetic acid 230 231 AA (Table 1, entry 11) was poorly reactive; the biobased levulinic acid (LA) gave 2c in good conversion (Table 1, entry 12), prompting us to include 40% of LA as a component of the GFS 232 instead of water, with the aim of using it both as peracid precursor and solvent component; however 233 234 in this case the epoxidation of cyclododecene 1b was not satisfactory (Table 1, entry 5). We also tested GFS-LA in combination with OA as peracid precursor on 1b; results were good but lower 235 236 than using GFS (Table 1, entries 3 and 4). The same happened with 1c (see SI, Table S1, entry 4). 237 Dimethyl carbonate DMC was also tested as peracid precursor but without good results (see SI, 238 Table S1, entries 1 and 3). The amount of the enzyme could be lowered till 30-25 U/mmol without 239 significant loss of reactivity (Table 1, 1b entries 7 and 8, 1c entry 14, 1d entries 18 and 19).

240 Limonene 1d is a very important biobased substrate, whose epoxy derivative is having some relevance in the field of polymer synthesis (Auriemma et al. 2015). Its internal double bond is much 241 more reactive than the terminal one, being electron-richer; in all the tested condition the product 2d 242 243 was obtained. Since we initially observed the formation of the diol as by-product (Table 1, entry 244 15), milder conditions were tested: i) halving the amount of NaDES; ii) keeping the temperature below 25 °C; iii) lowering addition rate of H₂O₂. All these conditions allowed to avoid the diol 245 formation (Table 1, entries 17 and 18). The use of a catalytic amount of acid precursor (Table 1, 246 entry 16) and a lower amount of the enzyme defined the best conditions to obtain 2d in very good 247 conversion (Table 1, entry 18). 248

As expected from limonene results, terminal bonds of styrene and itaconic anhydride were not reactive in the mild condition we tested for the other substrates (see SI, Table S2, entries 5 and 6).

251

252 *3.1.2 trans-Stilbene* **1***e*

253 *trans*-Stilbene **1e** is a challenging substrate because its double bond is electron-poor and it is poorly soluble in polar solvents like GFS. OA and other linear aliphatic carboxylic acids with shorter 254 (hexanoic HA, butanoic BA and acetic acid AA) and longer (dodecanoic acid, DA) chain lengths 255 were tested as peracid precursors (Table 2). In all cases OA resulted the most effective acid 256 257 precursor also in this case but 1 eq was needed to obtain an effective conversion (Table 2, entry 2). A decrease of the enzyme amount was possible, but a conversion of 75% was reached only after 48 258 259 h (Table 2, entry 4). Longer reaction times did not increase the conversion (SI, Table 2, entries 3 and 4). Differently, the electron-poor, α - β double bonds of crotonic acid and methyl crotonate were 260

very difficult to be epoxidized (see SI, Table S2, entries 7 and 8) and we obtained just traces of the
products. We also tested substrates carrying hydroxyl groups such as 1-octen-3-ol or *trans*-2-hexen1-ol but, as expected, the main product was the ester formed by OA and the alcohol under CAL-B
catalysis (data not shown).

265

266 *3.1.3 Oleic acid* **1***f*

Oleic acid **1f** is a very interesting substrate since its epoxide (9,10-epoxystearic acid) is a highly-267 valuable oleochemical due to its wide range of industrial applications, including cosmetics, personal 268 care, and pharmaceutical products. The epoxidation worked very well and without the addition of 269 270 OA (Scheme 2), thanks to an autocatalytic mechanism that formed the peroxy acid from the oleic acid itself. (Rüsch gen. Klaas & Warwel 1999) A temperature of 45 °C was required not only to 271 272 catalyze the reaction but also to avoid the product solidification. The condition used are the same 273 suggested and used by the recent literature (temperature at maximun 50 °C, an excess of H₂O₂, 274 short reaction time), except for the use of the solvent, which is generally toluene. (Milchert et al. 275 2015) The epoxidation can also be carried out in a solvent-free system, but the process is more 276 efficient for methyl oleate since the corresponding epoxide is liquid respect to solid 9,10epoxystearic acid. (Orellana-Coca et al. 2005) 277

278

279 <u>3.2 Baeyer-Villiger oxidations</u>

The first use of CAL-B as catalyst for B-V oxidations was performed in toluene with myristic acid 280 as peracid precursor. (Lemoult et al. 1995) Recent examples report ethyl acetate both as solvent and 281 282 peracid precursor (therefore in large excess with respect to the starting material) (Ríos et al. 2007; Rios et al. 2008; Chávez et al. 2013; Drożdż et al. 2013) and combination of ionic liquids and OA 283 284 as solvent and peracid precursor, respectively (OA in excess with respect to the starting ketone). 285 (Kotlewska et al. 2011; Drożdż et al. 2015) Urea-hydrogen peroxide is considered a milder oxidant than H_2O_2 alone and it was used to reduce the formation of water in the reaction, (Ríos et al. 2007; 286 287 Rios et al. 2008) nevertheless other studies showed no significant improvement in product 288 conversion and enzyme recycling. (Chávez et al. 2013) Considering that water is already present in 289 our GFS, the availability and the lower cost of hydrogen peroxide, this last one was thus chosen as 290 oxidant in our study. As for reaction times, when the reaction was carried out at room temperature it 291 generally required very long reaction times (in the order of days) to reach effective conversions. (Ríos et al. 2007; Rios et al. 2008; Chávez et al. 2013; Drożdż et al. 2013; Drożdż et al. 2015) 292

We tested B-V oxidation on various substrates (see section 2.4), using the same chemoenzymatic method in GFS previously described for epoxidation reactions: CAL-B, H_2O_2 (30% aqueous solution) and OA as peracid precursor (scheme in Table 3). Also in this case, a detailed study was conducted on the reaction conditions, with the aim of reducing the use of the reagents in excess and to use the mildest possible conditions.

298

299 *3.2.1 Cyclic ketones* **3a-c**

300 Since highly reactive in B-V oxidations, cyclohexanone 3a was the first substrate tested. By 301 carrying out the reaction at 20 °C, lactone 4a, (ε-caprolactone, Table 3, entry 1) was obtained but the reaction proceeded very slowly and an increase in time lead to the formation of the by-product. 302 6-hydroxyhexanoic acid 4a', caused by the ring-opening of 4a. Increasing the amount of catalyst or 303 304 temperature did not increase the selectivity towards 4a formation (Table 3, entries 2 and 3). Differently from the epoxidation reaction of cyclic alkenes (Table 1), the use of the peracid 305 306 precursor in catalytic amount did not give good results (Table 3, entry 4). Indeed, ω-hydroxy acid 307 formation is the main drawback in CAL-B mediated B-V oxidation (amounts reported in Table 3). 308 (X.-P. Wang et al. 2017) Increasing the amount of H₂O₂ to 2 eq (Table 3, entries 5 and 6) gave 309 higher conversions, without the formation of any by-product, while a shorter reaction time was achieved by conducting the reaction at 45 °C (Table 3, entry 6). The use of both OA and H₂O₂ in 310 excess (Table 3, entries 7 and 8) gave the best conversion: 74% at 20 °C and 58% at 45 °C. As 311

312 previous observed when the reaction was conducted at 45 °C, the reaction must be stopped after a 313 few hours to avoid by-product formation (Table 3, entry 8). Further increasing of both oxidant and

- acid amounts was not effective (Table 3, entry 9).
- The expected higher reactivity of cyclopentanone **3b** prompted us to lower the enzyme amount but
- also tuning temperature, oxidant and OA amounts (Table 3, entries 11-13) the high reactivity of the
- 317 substrate caused a rapid formation of the by-product **3b**'. As expected from the literature, (Chávez 318 et al. 2013; Drożdż et al. 2013; Drożdż et al. 2015) substrates with larger rings, as cyclooctanone
- 319 **3c**, are unreactive in all the tested conditions (Table 3, entry 14).
 - 320

321 3.2.2 2-phenylacetophenone **3d**

322 When using 2-phenylacetophenone 3d the regioselectivity issue must be considered, due to the formation of two possible regioisomers 4d and 4d' (structure in Table 4 foot) caused by the 323 324 migration of the phenyl group instead of the benzyl one (favored). As expected, we predominantly 325 obtained regioisomer 4d in 50% conversion at long reaction times (Table 4, entry 1). Higher 326 temperature did not increase the conversion but significantly increased the reaction rate (Table 4, 327 entries 1 and 3). Conversions decreased by lowering the amount of OA and enzyme (Table 4, entries 2 and 4). Using 2 eq of OA and H₂O₂ was not effective (Table 4, entry 5), while a great 328 329 excess of H₂O₂ gave 60% of 4d (Table 4, entry 6). Linear ketones and levulinic acid were tested but 330 the reactin did not work under the developed conditions (data not shown).

331

332 Conclusions

333 We demonstrated that chemoenzymatic oxidations using lipase CAL-B to form the active oxidant from carboxylic acid/H2O2 pair can be performed in a sugar-based NaDES composed by an 334 335 equimolar mixture of glucose, fructose, sucrose and water (GFS). Specific conditions to perform the 336 reaction on selected substrates in good conversion and selectivity were found. The best conditions for epoxidations proved to be related to the substrate reactivity; reaction conditions were tuned and 337 338 catalysts amounts decreased to obtain epoxides from poorly reactive and steric-hindered double 339 bonds (as *trans*-stilbene) and to control the formation of byproducts in more reactive alkenes (like 340 internal double bond of R-limonene). Baeyer-Villiger oxidations always required at least 341 stoichiometric amount of the peracid precursor to proceed and an excess of both oxidant and acid to 342 obtain good conversions. 343

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350 **Disclosure of interest**

- 351 The authors report no conflict of interest.
- 352
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Tables

Table 1. Epoxidation of cyclic alkenes with chemoenzymatic method in NaDES.

		$R_1 R_2$	CA Peracid p H ₂ O ₂ (Na 2 45 2	L-B orecursor 1 eq.) DES °C 0 h	$R_1 O R_2$		
Entry	Alkene	NaDES	CAL-B (U/mmol) Peracid precursor (eq)		Product conversion (%) ^a	By-products conversion (%) ^a	
	la				O 2a		
1	1a	ChCl- Sorb [1:1]	250	OA, 1	6	34 59	
	1b					-	
2	1b	ChCl- Sorb [1:1]	250	OA, 1	68		
3	1b	GFS	250	OA, 1	71		
4	1b	GFS [1:1:1] -LA	250	OA, 1	64		
5	1b	GFS [1:1:1] -LA	250	LA	15		
6	1b	GFS	250	OA, 0.1	75		
7	1b	GFS	25	OA, 1	80		
8	1b	GFS	25	OA, 0.1	79 (77)		
						-	
9	1c	GFS	250	OA, 1	>99		
10	1c	GFS	250	BA, 1	99		
11	1c	GFS	250	AA, 1	48		
12	1c	GFS	250	LA, 1	87		
13	1c	GFS	250	OA, 0.1	93		
14	1c	GFS	25	OA, 0.1	95 (91)		
					→	ОНОН	

	1d				2d	2d'
15 ^c	1d	GFS	60	OA, 1	76	14
16 ^c	1d	GFS	60	OA, 0.1	73	16
17 ^{c,d}	1d	GFS	60	OA, 1	89	-
18 ^{c,d}	1d	GFS	30	OA, 0.1	96	-
19°	1d	GFS	30	OA, 0.1	53	31

^a conversion by GC-MS, isolated yield in brackets; ^b diastereomeric ratio Z/E in **1b** and **2b** is always 2:1; ^c Room temperature (20 °C); ^d H₂O₂ total amount divided into 4 portions added in 4 hours. Acronyms: ChCl= choline chloride, Sorb = sorbitol, GFS = glucose, fructose, sucrose, H₂O (1:1:1:11), OA= octanoic acid, LA=

levulinic acid, AA = acetic acid, BA=butyric acid.

Table 2. Epoxidation of *trans*- stilbene 1e with chemoenzymatic method in GFS NaDES.

$1e \begin{array}{c} CAL-B \\ Peracid precursor \\ H_2O_2(1 \text{ eq.}) \\ \hline \\ GFS \\ 45 ^{\circ}C \\ 48 \text{ h} \end{array} \begin{array}{c} O \\ O \\ O \\ CAL-B \\ Peracid precursor \\ H_2O_2(1 \text{ eq.}) \\ \hline \\ GFS \\ 45 ^{\circ}C \\ 48 \text{ h} \end{array} \begin{array}{c} O \\ O $								
Entry	CAL-B (U/mmol)	Peracid precursor (eq)	2e conversion (%) ^a					
1 ^b	250	OA, 1	60					
2 ^b	250	OA, 0.1	traces					
3	250	OA, 1	74 (70)					
4	25	OA, 1	73					
5	25	DA, 1	11					
6	25	HA, 1	54					
7	25	BA, 1	-					
8	25	AA, 1	-					

^a conversion by GC-MS, isolated yield in brackets; ^b time (20h)

Acronyms: GFS = glucose, fructose, sucrose, water (1:1:1:11), OA= octanoic acid, DA = dodecanoic acid, HA= Hexanoic Acid, BA=butyric acid, AA = acetic acid.

			0 0	CAL-B Octanoic Act H ₂ O ₂	d	O	D	
		R ₁	[^] R ₂ ⁻	GFS T°C time		R ₁ O	ι π ₂	
Entry	Ketone	CAL-B (U/mmol)	OA (eq)	H ₂ O ₂ (eq)	Т (°С)	time (h)	Product conversion (%) ^a	By-product conversion (%)l ^a
	O J J Ja							ноон 4а'
1	3 a	125	1	1	20	20 70	32 23	- 53
2	3a	250	1	1	20	20	21	16
3	3 a	200	1	1	45	5 15	35 36	- 28
4	3 a	125	0,1	1	20	20 40	15 15	-
5	3 a	125	1	2	20	20 40	37 55	-
6	3 a	125	1	2	45	5 20	30 20	- 8
7	3 a	125	2	2	20	20 40 64	49 61 74	
8	3 a	125	2	2	45	5 20	52 29	- 21
9	3 a	125	1	3	20	20 4d	36 54	- 16
10	3 a	125	3	3	20	20 4d	58 54	- 16
								но
	30						4b	4b'
11	3b	65	1	1	20	20 40	15 7	- 50
12	3 b	65	1	1	45	20	10	30
13	3 b	65	0.1	1	20	40	4	15
								Not found
14	3c	65	various	various	20	various	traces	-

Table 3. Baeyer-Villiger oxidation of lactones with chemoenzymatic method in GFS.

^a conversion by GC-MS, isolated yield in brackets; Acronyms: GFS = glucose, fructose, sucrose, H₂O (1:1:1:11), OA= octanoic acid.

Table 4. Baeyer-Villiger oxidation of 2-phenylacetophenone **3d** in GFS

$\begin{array}{c c} & & CAL-B & & O \\ & & & Octanoic Acid \\ & & H_2O_2 \\ & & & GFS \\ & & 45^{\circ}C \\ & & time \end{array} \qquad $							
Entry	Ketone	CAL-B (U/mmol)	OA (eq)	H ₂ O ₂ (eq)	time (h)	Conversion 4d (%) ^{a,c}	
1 ^b	3d	250	1	1	40 50 7 days	7 17 50	
2 ^b	3d	250	0.1	1	50	12	
3	3d	250	1	1	40 7 days	40 (34) 50	
4	3d	100	1	1	20 5 days	18 27	
5	3d	250	2	2	20 50 3 days	38 42 44	
6	3d	250	1	3	23 4 days	55 60	

^a conversion by GC-MS, isolated yield in brackets; ^b temperature (20 °C); ^c in all the entries there are traces of the regiosomer of 4d, 4d', 4d'.

Acronyms: GFS = glucose, fructose, sucrose, $H_2O(1:1:1:11)$, OA = octanoic acid.

Schemes

Scheme 1



Schemes Captions

Scheme 1. Chemoenzymatic pathway for epoxidations and Baeyer-Villiger oxidations. Scheme 2. Epoxidation of oleic acid with chemoenzymatic method in GFS.