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¹ Use of Monosaccharides in Metal-Catalyzed Coupling Reactions

² Sara H. Kyne^{*,†} and Jason E. Camp^{*,‡}

3 [†]School of Chemistry, University of Lincoln, Lincoln LN6 7TS, United Kingdom

4 [‡]Department of Chemical Sciences, University of Huddersfield, Huddersfield HD1 3DH, United Kingdom

5 ABSTRACT: The addition of monosaccharides to metal-6 catalyzed coupling reactions can be beneficial in terms of decreasing the time required, chemical waste products and 7 overall cost of the process. Monosaccharides are used in a 8 number of different ways, including (a) acting as a ligand for 9 the metal, (b) providing the appropriate reduction potential 10 for a chemical process and (c) acting as a reducing agent for 11 the formation and stabilization of catalytically active metal 12 nanoparticles. Recently, there has been a significant amount of 13 research in this growing field and there is thus the potential for 14 the addition of monosaccharides to coupling reactions to have 15 a significant impact on the synthesis of the important small 16 molecules on which we have all come to rely. This Perspectives 17



- 18 Article will cover recent developments in the addition of monosaccharides to metal-catalyzed coupling reactions with an 19 emphasis on their utility and limitations in order to facilitate the further development of this exciting area of research.
- 20 KEYWORDS: Monosaccharide, Biorenewables, Metal catalysis, Green chemistry, Nanoparticle

21 INTRODUCTION

22 Metal-catalyzed coupling processes are a ubiquitous part of the 23 modern chemists' toolkit for the synthesis of added-value small 24 molecules on which we have all come to rely. To make these 25 processes more efficient, in terms of time, expense and cost to 26 the environment, unmodified monosaccharides have been 27 added to metal-catalyzed reactions as part of research into 28 the use of biorenewables in catalytic/chemical reactions. The 29 addition of monosaccharides can serve many purposes in these 30 reactions, including (a) acting as a ligand for the metal, (b) 31 providing the reduction potential for a chemical process and (c) 32 acting as a reducing agent for the formation and stabilization of 33 catalytically active metal nanoparticles. The ability of 34 monosaccharides to reduce metals has been known for 35 decades,¹ for example Benedict's² or Fehling's tests,³ but their 36 use in cross-coupling reactions has flourished in recent years. 37 This review will focus on the latest uses of monosaccharides in 38 metal-catalyzed coupling reactions. Because of recent reviews 39 and full publications, the following areas will not be covered in 40 this review: polysaccharides,⁴⁻⁹ smaller sugar derived alde-41 hydes/carboxylic acids (Leuckart–Wallach reaction) $^{10-16}$ or 42 reactions in which sugars are used as starting materials or 43 incorporated into the molecule.^{17–24}

44 MONOSACCHARIDES AS LIGANDS

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45 One of the most common uses of monosaccharides in metal-46 catalyzed reactions are as ligands for a catalytically active metal 47 species.²⁵ For example, Sekar and Thakur recently disclosed the 48 synthesis of phenols **2** from aryl halides **1** in a process that was 49 catalyzed by a copper/glucose system (Scheme 1).²⁶ Aryl 50 iodides and bromides **1** were reacted with excess potassium

Scheme 1. Synthesis of Phenols 2 from Aryl Halides 1



hydroxide (4–8 equiv) in the presence of copper(II) acetate (5 $_{51}$ mol %) and D-glucose (5 mol %) to give good to excellent $_{52}$ yields of the corresponding phenols **2**. The reactivity of aryl $_{53}$ chlorides depended on the nature of the electron withdrawing $_{54}$ group with substrates containing a nitro group giving an $_{55}$ excellent yield of phenol **2**.

Recently, a number of carbon–nitrogen cross-coupling 57 reactions have been developed employing a catalytic system 58 formed *in situ* from copper(I) iodide and D-glucosamine in the 59 presence of base.²⁷ For example, anilines were formed from aryl 60 halides in the presence of excess aqueous ammonia (10 equiv) 61 or sodium azide (3 equiv).^{28,29} Zhang et al. reported the use of 62 similar conditions for the cross-coupling of aryl halides 3 with 63 nitrogen heterocycles 4 (1.2 equiv.; Scheme 2).³⁰ Most of the 64 s2 examples used imidazole as the heterocycle 4, and good yields 65 were observed for aryl iodides 3 bearing electron-withdrawing 66

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Scheme 2. Cross-Coupling of Aryl Halides 3 with Nitrogen Heterocycles 4



67 groups. Unfortunately, the reaction did not occur with aryl 68 chlorides.

⁶⁹ Zhang et al. extended this methodology to carbon–sulfur ⁷⁰ cross-coupling reactions. In this work, aryl iodides **6** were ⁷¹ reacted with diphenyl disulfide (7, 0.6 equiv) in the presence of ⁷² copper(I) iodide (10 mol %), D-glucosamine (10 mol %) and ⁷³ cesium carbonate (2 equiv) to give the corresponding ⁷⁴ unsymmetrical diaryl sulfide **8** (Scheme 3).³⁰ When aryl





75 bromides were tested the reaction occurred, but required 24 76 h to go to completion. Similar methodology was used by the 77 same group to synthesize a variety of diaryl sulfones from aryl 78 halides and sodium benzenesulfonates.³¹

79 D-Glucosamine has also been successfully employed as a 80 ligand in iron-catalyzed Grignard cross-coupling reactions of 81 vinylic **10** and allylic bromides **11** (Scheme 4). Phenyl- or

Scheme 4. D-Glucosamine Successfully Employed as a Ligand in Iron-Catalyzed Grignard Cross-Coupling Reactions



⁸² benzylmagnesium bromides **9** were reacted with bromides **10** ⁸³ or **11** in the presence of iron(II) acetylacetonate (5 mol %) and ⁸⁴ D-glucosamine hydrochloride (5 mol %). Triethylamine (5 mol ⁸⁵ %) was added to deprotonate the ligand and thus increase its ⁸⁶ solubility in THF.³² Moderate yields of substituted alkenes **12** ⁸⁷ were obtained from allylic bromides **10**, and good yields of the ⁸⁸ sp³-hybridized products **13** were obtained from alkenyl ⁸⁹ bromides **11**.

⁹⁰ D-Glucosamine was shown to improve the yield in palladium-⁹¹ catalyzed Mizoroki–Heck reactions of aryl halides (Scheme ⁹² 5).³³ Aryl halides **14** were reacted with activated alkenes **15** ⁹³ (1.2 equiv) in the presence of palladium(II) acetate (0.5 mol ⁹⁴ %), D-glucosamine (1 mol %) and potassium carbonate (2.0 ⁹⁵ equiv). Aryl iodides and bromides afforded good to excellent ⁹⁶ conversion to stilbenes **16** with unsubstituted and *para*-⁹⁷ substituted electron withdrawing groups. Conversion was Scheme 5. D-Glucosamine Shown To Improve Yield in Palladium-Catalyzed Mizoroki–Heck Reactions of Aryl



moderate when the arene was substituted at the *ortho* position. 98 Aryl chlorides 14 reacted, albeit with low conversion (5–25%). 99

MONOSACCHARIDES FOR NANOPARTICLE FORMATION

One of the most common uses of monosaccharides in organic 102 transformations are as reductants for the formation of metal 103 nanoparticles, in which the sugar serves to reduce the metal in 104 the presence of a template.³⁴⁻⁴⁵ Monosaccharides, glucose in 105 particular, have also been used as supports for metal 106 nanoparticles.⁴⁶⁻⁴⁹ In some cases, the monosaccharides act as 107 both the stabilizer for the metal nanoparticles as well as the 108 reducing agent.⁵⁰⁻⁵⁴ Alternatively, additional reducing agents 109 can be added to the mixture of sugar and metal if required. For 110 example, monodispersed colloidal carbon spheres have been 111 synthesized by a two-step hydrothermal approach under mild 112 conditions by Sun et al.55 In this work, separating the 113 nucleation and growth steps allowed for a narrow size 114 distribution with diameters ranging from 160 to 400 nm. 115 Interestingly, the size distribution decreased with an increasing 116 concentration of glucose. D-Glucose has also been used as the 117 metal nanoparticle support. In this case, palladium(0) nano- 118 particles were synthesized by the reduction of $H_2[PdCl_4]$ or 119 $[Pd(NH_3)_4Cl_2]Cl_2$ in the presence of excess hydroxylamine and 120 D-glucose under ambient, aqueous conditions.⁵⁶ Character- 121 ization of the palladium nanoparticles revealed magnetization 122 differences depending on the oxidation state of the palladium 123 precursor. TEM analysis revealed that when starting from the 124 Pd(II) complex, the nanoparticles were an average size of 6 nm 125 and polydispersed, whereas starting from the Pd(IV) complex 126 formed nanoparticles with an average size of 8 nm that were 127 mainly monodispersed. In 2004, Sun and Li reported the 128 synthesis of colloidal carbon spheres starting from glucose, 129 which underwent subsequent functionalization due to the 130 reactive surface present.⁵⁷ For example, the FTIR spectrum 131 revealed the existence of carbonyl and hydroxyl groups that 132 maintained the hydrophilicity of the carbon spheres. Colloidal 133 carbon spheres were prepared from aqueous glucose by 134 hydrothermal synthesis, undergoing aromatization and carbon- 135 ization to form 200 nm carbon spheres at 160 °C in 3.5 h, and 136 1500 nm at 180 °C in 10 h. Under reflux, palladium(0) 137 nanoparticles were loaded onto the surface, covering the carbon 138 spheres with a uniform shell of 10-20 nm palladium. In related 139 methodology, Zhang et al. described the preparation of highly 140 dispersed, narrow diameter palladium nanoparticles on carbon 141 spheres via in situ reduction.⁵⁸ Precise control of the dispersity 142 and size of the palladium(0) nanoparticles was possible by 143 careful adjustment of the reaction conditions (temperature, 144 time, pH and ratio of palladium(0) to carbon spheres). 145 Homogeneously distributed, small diameter (7.7 nm) palladium 146 crystals were prepared on carbon spheres at pH 7.0 in ethanol 147 at 70 °C. 148

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The isolated metal nanoparticles have subsequently been used in a number of important catalytic processes. For example, so used in a number of important catalytic processes. For example, so used in a number of important catalytic processes. For example, so used in a number of important catalytic processes. For example, so used in a number of important catalytic processes. For example, so used in a number of important catalytic processes. For example, so used in a number of important catalytic processes. For example, so used in a number of important catalytic processes. For example, so used in a number of important catalytic processes. For example, so used in a number of important catalytic processes. For example, so used in a number of important catalytic processes. For example, so used in a number of important catalytic processes. For example, so used in a number of important catalytic processes. For example, so used in a number of important catalytic processes. For example, so used in a number of important catalytic processes. For example, so used in a number of important catalytic processes. For example, so used in a number of important catalytic processes. For example, so used in a number of important catalytic processes. For example, so used in a number of important catalytic processes. For example, so used in a number of important catalytic processes. For example, so used in a number of important catalytic processes. For example, so used in a number of important catalytic processes. For example, so used in a number of important catalytic processes. For example, so used in a number of important catalytic processes. For example, so used in a number of important catalytic processes. For example, so used in a number of important catalytic processes. For example, so used in a number of important catalytic processes. For example, so used in a number of important catalytic processes. For example, so used in a number of important catalytic processes. For example, so used in a number of important catalytic process

Scheme 6. Palladium(0) Nanoparticle Catalyzed Domino Sonogashira-Cyclization Reaction to Synthesize Various Isoindolinones 20 and Furoquinolines 21



155 by the procedure of Sarkar et al. in which $H_2[PdCl_4]$ was 156 reduced in the presence of hydroxylamine and D-glucose under 157 aqueous conditions.⁵⁶ Following the reaction, the catalyst could 158 be recovered in high yield and a recycling study showed only a 159 gradual decrease in activity for up to five subsequent reactions 160 before significant loss of yield was observed.

MONOSACCHARIDES AS REDUCTANTS IN CHEMICAL PROCESSES

163 Monosaccharides can also be added to organic transformations 164 to act as reductants for chemical process that occur in the 165 absence of catalyst^{60,61} or to reduce a metal precatalyst to the 166 necessary oxidation state *in situ* so that the reaction can 167 proceed. This concept has been employed for a variety of 168 reactions such as dehalogenations,⁶² reductions^{63–66} and 169 coupling processes. Glucose can also be used for the *in situ* 170 formation of the active metal catalyst. For example, Cuevas-171 Yañez et al. showed that addition of 25 mol % of glucose to the 172 reaction mixture resulted in an increased yield of the desired 173 triazoles **24** from alkynes **22** and azides **23** under copper-174 catalyzed click reaction conditions (Scheme 7).⁶⁷

Scheme 7. Increased Yield of the Desired Triazoles 24 from Alkynes 22 and Azides 23 under Copper-Catalyzed Click Reaction Conditions



In related work, Singh et al. showed that it was beneficial to add glucose to copper-catalyzed click reactions⁶⁸ that were part multicomponent coupling reactions under microwave conditions (Scheme 8). Thus, the three component reaction phenylazides **25**, 4-(prop-2-yn-1-yloxy)benzaldehydes **26**, and 1,2-diaminobenzenes **27** afforded the triazole adducts **28** in good yields (Scheme 8a).⁶⁹ Additionally, a four component process resulted in the efficient formation of 3-phenyl-2-[4-{(1-

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phenyl-1*H*-1,2,3-triazol-4-yl)methoxy}phenyl]thiazolidin-4-183 ones **29** from readily available starting materials (Scheme 8b).⁷⁰ 184 In both cases, the glucose is purported to reduce the copper to 185 the catalytically active species. Related work by Wan et al. 186 demonstrated that copper—glucose systems catalyzed the three 187 component reactions of phenols, acyl chlorides and Wittig 188 reagents to form β -aryloxy acrylates.⁷¹ Furthermore, Guchhait 189 et al. developed a novel A³-coupling methods for the synthesis 190 of *N*-fused imidazoles using a copper(II) sulfate—glucose 191 catalyst.⁷² This methodology was subsequently harnessed by 192 Iyer et al. for the synthesis of luminescent imidazo[1,2-193 *a*]pyridines.⁷³

MONOSACCHARIDES FOR THE IN SITU 195 FORMATION/STABILIZATION OF CATALYSTS 196

One of the most important advances in this area is the ability to 197 form catalytically active metal nanoparticles in situ from 198 unmodified reducing sugars and subsequently recycle the 199 catalyst. Traditionally, bioderived metal catalysts need to be 200 synthesized and isolated prior to reaction, and are frequently 201 difficult to recycle following the reaction, which can increase 202 the amount of time, chemical waste and expense of the overall 203 process.⁷⁴⁻⁸¹ Building upon the work in nanoparticle formation 204 from reducing sugars, and the use of reducing sugars as 205 reductants in catalytic processes, Nacci et al. recently disclosed 206 an Ullman type homocoupling of aryl halides catalyzed by in 207 situ generated palladium(0) nanoparticles (Scheme 9).⁸² Thus, 208 s9 the homocoupling of bromo- and chloroarenes 30 in the 209 presence of glucose (0.5 equiv), palladium(II) acetate (3 mol 210 %) and tetrabutylammonium hydroxide (3.0 equiv) afforded 211 the desired biaryls 31 in good yield. In this process, the glucose 212 is postulated to both reduce the palladium(II) acetate to the 213 catalytically active palladium(0) species, as well as stabilize the 214 in situ formed catalyst through the formation of nanoparticles. 215 In contrast to other related reports (vide infra), exogenous 216 capping agents were used in this study. TEM analysis was used 217 to confirm the formation of nanoparticles, which had an 218 average particle size around 15 nm, and XPS was used to show 219 that the palladium in the isolated nanoparticles was in the zero 220 oxidation state. A recycling study demonstrated that the 221 catalytic solution was active for 3 cycles, but the yield decreased 222 precipitously thereafter. 223

Nacci et al. also showed that an Ullman type homocoupling 224 of haloarenes 32 to give the corresponding biaryls 33 could be 225 facilitated by gold nanoparticles formed in situ from the 226 reduction of gold(III) acetate (2 mol %) with a stoichiometric 227 amount of glucose (Scheme 10).83 This catalytic system was 228 s10 active for bromo- and iodo-substituted arenes 32, as well as 229 alkenes. The less reactive aryl chlorides were found to be 230 unreactive under the reaction conditions. It was also found that 231 the ionic liquid tetrabutylammonium acetate (TBAA) was a 232 competent solvent and base for the reaction, which could 233 substitute for water. Interestingly, the nanoparticles formed in 234 the ionic liquid were much larger than those formed in water 235 (ca. 2 nm vs. Twenty nm respectively) and in general resulted 236 in a decreased yield compared to the aqueous conditions. 237 Unfortunately, attempts at recycling these catalysts showed a 238 similar poor performance to the palladium system discussed 239 above (cf. Scheme 9). 240

Recently, we disclosed methods for the use of glucose 241 derived palladium(0) nanoparticles as *in situ* formed catalysts 242 for Suzuki–Miyaura cross-coupling reactions in the green 243 solvent isopropyl alcohol (Scheme 11).⁸⁴ The cross-coupling of 244 s11 Scheme 8. Four Component Process Resulted in the Efficient Formation of 3-Phenyl-2-[4-{(1-phenyl-1*H*-1,2,3-triazol-4-yl)methoxy}phenyl]thiazolidin-4-ones 29



Scheme 9. Ullman Type Homocoupling of Aryl Halides Catalyzed by *in Ditu* Generated Palladium(0) Nanoparticles



Scheme 10. Ullman Type Homocoupling of Haloarenes 32 To Give Corresponding Biaryls 33 Facilitated by Gold Nanoparticles Formed *in Situ* from Reduction of Gold(III) Acetate with Stoichiometric Amount of Glucose



Scheme 11. Use of Glucose Derived Palladium(0) Nanoparticles as *in Situ* Formed Catalysts for Suzuki– Miyaura Cross-Coupling Reactions Green Solvent Isopropyl Alcohol



245 aryl iodides **34** and aryl boronic acids **35** in the presence of 246 palladium(II) acetate (1 mol %) and glucose (5 mol %) gave 247 the desired biaryls **36** in moderate to good yields under either 248 thermal or microwave heating conditions. In contrast to the 249 reports of Nacci et al., only a small amount of glucose was 250 required and no capping agents were employed. EF-TEM analysis of the *in situ* formed nanoparticles showed that the 251 palladium was surrounded by a hydrophilic, hydroxylated shell. 252 The hydrophilic/polar nature of the nanoparticles allowed for 253 their facile removal from the cross-coupled product. ICP-MS 254 analysis showed a 65% decrease in the amount of metal 255 incorporated into the final compounds compared to reactions 256 that did not contain glucose. Interestingly, Jiang and Fossey et 257 al. have found that monosaccharides bind to boronic acid to 258 form the less reactive boronate ester. They used the retardation 259 of the Suzuki–Miyaura homocoupling reaction to develop 260 fluorescent sensors for glucose detection.^{85,86}

Subsequently, Jain et al. described the use of reducing sugars 262 in palladium mediated cross-coupling reactions, in which the 263 metal was catalyzing multiple, mechanistically distinct steps; 264 autotandem catalysis^{87,88} (Scheme 12).⁸⁹ After screening nine 265 s12

Scheme 12. Use of Reducing Sugars in Palladium Mediated Cross-Coupling Reactions



difference reducing sugars, they found that the addition of 266 mannose (3 equiv) gave the desired cross-coupled products of 267 Suzuki–Miyaura and Mizoroki–Heck reactions, while con- 268 currently reducing the nitro functionality to an aniline. For 269 example, reaction of halo-nitrobenzenes **37** with arylboronic 270 acids **38** in aqueous DMF at 130 °C (microwave) gave the 271 coupled biaryl anilines **39** in moderate to excellent yields. 272 Similarly, the reaction of iodo-nitrobenzenes **40** with styrenes 273 **41** under the same reaction conditions afforded substituted 274 amino-stilbenes **42** in good to excellent yields. It is unclear from 275 the analysis conducted by the researchers whether the mannose 276 is simply acting as a ligand and source of hydrogen under the 277 reaction conditions, or if it is also stabilizing *in situ* formed 278 nanoparticles.

Building upon our work discussed above, we recently 280 reported the use of glucose derived nanoparticles for the 281 282 Mizoroki–Heck, Sonogashira and Suzuki–Miyaura cross-283 coupling reactions in aqueous solvents (Scheme 13).⁹⁰ The

Scheme 13. Use of Glucose Derived Nanoparticles for the Mizoroki-Heck, Sonogashira and Suzuki-Miyaura Cross-Coupling Reactions in Aqueous Solvents



284 reaction of aryl halides 43 with alkenes 44 or alkynes 46 285 proceeded in moderate to excellent yields to afford stilbenes 45 286 or substituted alkynes 47, respectively. The palladium(0)287 nanoparticle catalysts were formed in situ from palladium(II) acetate (2 mol %) via the addition of glucose (4–10 mol %) to 288 the reaction. In addition, a Suzuki-Miyaura protocol for the 289 290 synthesis of biaryls 50 in aqueous DMF was developed using 291 the same in situ derived palladium(0) nanoparticles. Impor-292 tantly, this protocol was also viable for arvl bromides. In contrast to the study by Jian et al.,⁷⁴ the nitro functionality was 293 294 not reduced in any of the three cross-coupling reactions that 295 were investigated. This is possibly due to the relatively small 296 amount of glucose that was added to the reaction (cf. Scheme 297 12). EF-TEM analysis of the in situ formed nanoparticles 298 confirmed that the palladium was surrounded by a hydrophilic, 299 hydroxylated shell. The nature of this shell allowed for the facile 300 partitioning of the catalyst between the aqueous and organic 301 phases, which enabled catalyst recycling for up to four cycles 302 without significant loss of activity.

303 CONCLUSION

304 The importance of the addition of monosaccharides to metal-305 catalyzed processes in organic chemistry has expanded rapidly 306 in recent years. These biorenewable materials can be used for a 307 number of important processes including as ligands for a metal 308 catalyst, to provide the appropriate reduction potential for a chemical process, and as a reducing agent for the formation and 309 310 stabilization of catalytically active metal nanoparticles. These 311 recent developments in the field will provide the basis for 312 further rapid advancements. Looking forward, catalytic 313 processes in which the reducing potential of renewable sugars is harnessed for the generation, stabilization and turnover of 314 315 catalytically active metal nanoparticles, sugar-powered catalysis, 316 will be developed. These processes have the potential to make 317 existing protocols greener in terms of time, expense and cost to 318 the environment, as well as allowing for the development of 319 novel metal-catalyzed processes that are currently not possible. 320 Additionally, the inherent chirality of the monosaccharides will 321 be harnessed in order to develop catalytic access to 322 enantiomerically enriched products. In conclusion, the addition 323 of monosaccharides to metal-catalyzed processes has resulted in 324 a number of important new methods that allow access to the

small molecules on which we have all come to rely. It is 325 expected that innovative new applications will be developed 326 that build on this exciting research. 327

AUTHOR INFORMATION	328
Corresponding Authors	329
*S. H. Kyne. Email: skyne@lincoln.ac.uk. Tel: +44 (0)15 2283	330
7402.	331
*J. E. Camp. Email: j.e.Camp@hud.ac.uk. Tel: +44 (0)14 8447	332
3180.	333
Notes	334
The authors declare no competing financial interest.	335
Biographies	336



Sara H. Kyne obtained her Ph.D. from The University of Melbourne 337 (Australia) under the supervision of Prof. Carl H. Schiesser in the area 338 of physical—organic intramolecular radical chemistry. She then 339 undertook a postdoctoral position with Prof. Jonathan M. Percy 340 (University of Strathclyde, UK) and a Marie Curie Intra European 341 Fellowship with Prof. Louis Fensterbank (Université Pierre et Marie 342 Curie, France). She is currently a lecturer at the University of Lincoln 343 working on the development of sustainable catalysis and radical 344 methodology. 345



Jason E. Camp obtained his Ph.D. from The Pennsylvania State 346 University under the supervision of Prof. Steven Weinreb working on 347 the total synthesis of the chartellamide and chartelline family of marine 348 natural products. He then was a postdoctoral fellow working with Prof. 349 Donald Craig (Imperial College London) before obtaining lecture- 350 ships at the University of Nottingham and Queen Mary University of 351 London. He is currently a senior lecturer at the University of 352 Huddersfield working on the development of novel autotandem 353 catalysis methods as well as sugar-powered catalysis protocols. 354

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