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### Spotlight

# Sodium Persulfate (Na<sub>2</sub>S<sub>2</sub>O<sub>8</sub>)

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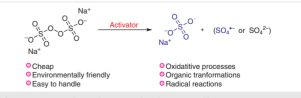
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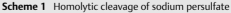
**Key words** oxidation, persulfates, radical reaction, sulfate radical anions, inorganic reagents

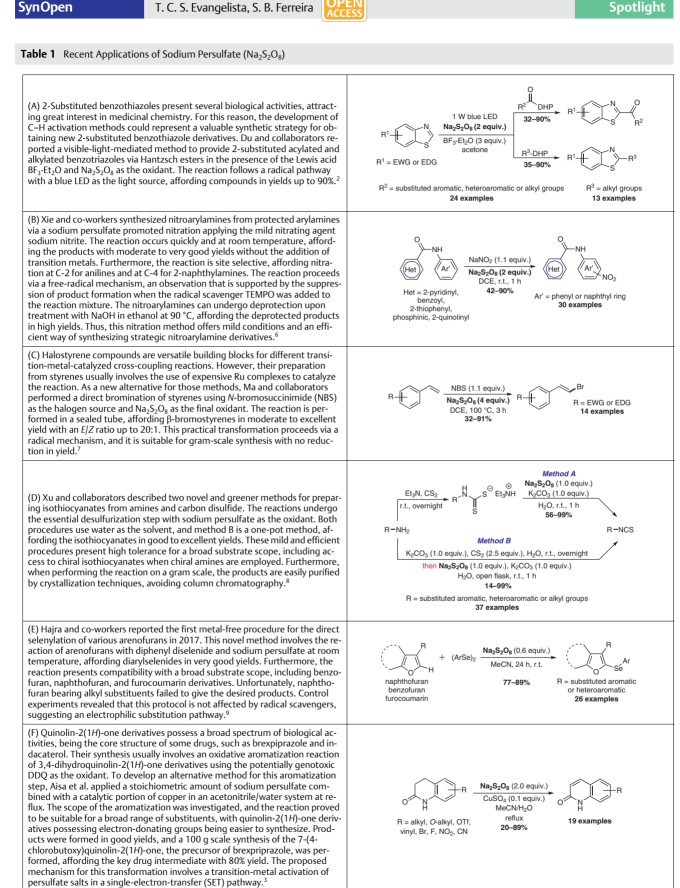
Sodium persulfate, or sodium peroxodisulfate, is an environmentally friendly inorganic compound commonly used as an oxidizing agent in chemistry. This reagent undergoes homolytic cleavage in solutions, affording sulfate radical anions that can present several applications (Scheme 1). Recently,  $SO_4$  has been applied in diverse processes, such as the degradation of antibiotics and dyes in wastewater treatment.<sup>1</sup> In organic synthesis, sodium persulfate can be used as an oxidant in transition-metal-catalyzed reactions or as an oxidative species in crucial steps in metal-free reactions.<sup>2,3</sup> Na<sub>2</sub>S<sub>2</sub>O<sub>8</sub> is an inexpensive oxidant that is stable and easy to handle, making it a good reagent choice for several strategic synthetic transformations.<sup>4</sup> It is synthesized industrially through an electrolytic oxidation process from sodium hydrogen sulfate,<sup>5</sup> and recent applications for this compound are presented herein (Table 1).



Tereza Cristina Santos Evangelista received her MSc in chemistry from the Federal University of Rio de Janeiro in 2018. Currently, she is a PhD student at the same institution under the supervision of Prof. Sabrina B. Ferreira and Prof. Carlos R. Kaiser. Her work involves the synthesis and biological evaluation of novel nitrogenated heterocycles. **Sabrina Baptista Ferreira** received her PhD from the Federal University of Rio de Janeiro (UFRJ) in 2008 under the supervision of Prof. Carlos R. Kaiser and Prof. Vitor F. Ferreira. After the postdoctoral experience held at Fluminense Federal University (UFF) with Prof. Vitor F. Ferreira, she became a professor at UFRJ in 2010 where she is the head of the Laboratory of Organic Synthesis and Biological Prospecting. Her research efforts focus on the organic synthesis, acting on the following subjects: heterocycles, carbohydrates, nucleosides and naphthoquinones, search for biologically active compounds.







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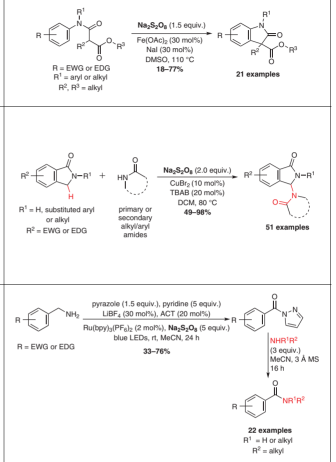
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### Spotlight

(G) To expand the scope of application for the oxidative system Fe(OAC)<sub>2</sub>/Nal/Na<sub>2</sub>S<sub>2</sub>O<sub>8</sub>, Zhou and co-workers carried out an intramolecular dehydrogenative coupling reaction to afford pharmacologically relevant oxindazole derivatives from a variety of anilides. For the reaction optimization, different oxidants were tested, including K<sub>2</sub>S<sub>2</sub>O<sub>8</sub>, (NH<sub>4</sub>)<sub>2</sub>S<sub>2</sub>O<sub>8</sub>, and Na<sub>2</sub>S<sub>2</sub>O<sub>8</sub>, where the latter gave the best results. The scope and feasibility of the proposed method were screened by reacting a series of anilide derivatives. The reaction presented broad substrate scope, and electron-rich aromatic derivatives gave the best product yields. The reaction swithout a drop in yield.<sup>10</sup>

(H) The amide functional group has a central importance in biologically active compounds. For this reason, straightforward methods to prepare this functionality are very desirable. Inspired by the cross-dehydrogenative-coupling (CDC) strategy, Zha and collaborators described the direct amidation of isoindolinone derivatives using sodium persulfate as the oxidant. The reaction optimization was performed using methylisoindolin-1-one and pyrrolidin-2-one as model substrates. To improve the formation of  $SO_4$  - species, catalytic amounts of transition metals were applied, including CuBr<sub>2</sub>, Pd(OAc)<sub>2</sub>, AgNO<sub>3</sub>, and FeCl<sub>2</sub>, where CuBr<sub>2</sub> gave the best results. The optimized procedure was applied to the synthesis of a variety of amide derivatives, possessing a broad substrate scope and great selectivity towards obtaining the amide functionality primarily at the C3 position of the isoindolinone motifs. The application of the method was extended to the functionalization of indobufen with different amides, which can be applied in structure-activity studies in drug design and to late-stage drug functionalization. Furthermore, control experiments suggested that the transformation involves a radical and electrophilic pathway

(I) In connection with photoamidation strategies, Leadbeater and co-workers reported for the first time the use of a tandem oxidative pathway for the conversion of amines into amides. The proposed process occurs via a visible-light-promoted reaction with the dual catalyst  $Ru(bpy)_3(PF_6)_2$  and 4-ACNH-TEMPO in the presence of the terminal oxidant sodium persulfate. This process was investigated for the synthesis of *N*-acylpyrazoles, applying a blue LED as the light source. Control experiments were performed, proving that the photocatalyst, the light source, and catalyst system are vital for the product formation. Furthermore, absence of  $Na_2S_2O_8$  led to the formation of the homocoupled imine. The authors also investigated the reaction scope for the tandem process followed by a transamination reaction. The two-step method occurs with no need for purification of the *N*-acyl pyrazole intermediate. This protocol, followed by the transamination reaction also proved to be broad in substrate scope, affording a variety of amines with moderate to good yields.<sup>12</sup>



## **Conflict of Interest**

The authors declare no conflict of interest.

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