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Photonic contacting of gas–liquid phases in a falling film microreactor for continuous-flow photochemical catalysis with visible light†

Thomas H. Rehm,^{*a} Sylvain Gros,^{ab} Patrick Löb^a and Albert Renken^b

A microstructured falling film reactor was applied to the dye-sensitized photochemical conversion of 1,5-dihydroxynaphthalene to juglone. This continuous-flow microreactor enables the efficient contacting of a gas and a liquid phase in combination with external irradiation by high-power LED arrays offering various wavelengths. Two sensitizers were used for the photochemical *in situ* generation of singlet oxygen as key step in the synthesis of the natural product juglone. The photochemical process was investigated according LED wavelength, LED power, oxygen partial pressure, reactor architecture, substrate concentration and flow rate, and optimized to a conversion of $X = 97\%$ with 99% selectivity. Based on the experimental results process parameters like quantum efficiency, productivity and space time yield were calculated and used for the evaluation of the photochemically catalyzed synthesis of juglone in continuous-flow mode.

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Introduction

Photochemically catalysed reactions have become of fundamental interest in the recent years.^{1–5} Especially with the advent of photocatalytically active compounds and materials, which are able to absorb light in the visible region of the electromagnetic spectrum, many new synthetic methods have been developed for the preparation of valuable fine chemicals under mild reaction conditions.^{6–15} A broad variety of transformations can be performed with organometallic compounds like Ru- or Ir-based polypyridyl complexes.^{16,17} In conjunction with organocatalysts as chiral inductors stereoselective synthesis has been made possible as well.^{18–22} More recently the application of precious metal-free catalysts has amplified the development of cost-efficient synthetic protocols for photochemical reactions.^{23–26} Meanwhile it has been generally accepted that light, *i.e.* photons with defined wavelength, is a highly potent, selective and, in terms of *Green Chemistry*, a traceless reagent for synthetic organic chemistry.^{27,28}

The dye-sensitized activation of molecular oxygen offers an important strategy to incorporate oxygen into molecules

under mild reaction conditions.²⁹ A first industrial application of photochemical oxygen activation has made its way into production. The *Sanofi* process for the production of artemisinin is one important example for the *in situ* formation of singlet oxygen using tetraphenylporphyrine as visible light-absorbing sensitizer.³⁰ Starting in 2013 the final production scale of 60 tons per year has already been reached in 2014. Two important prerequisites must be fulfilled in this case for a successful realization of dye-sensitized singlet oxygen formation: efficient oxygen gas contacting with the liquid phase and thorough irradiation of the reaction composition. In general, photochemical batch equipment, *e.g.* glass flasks and immersion-well setups with pressurized mercury lamps, often lack of efficient gas–liquid contacting and uniform irradiation of the reaction solution, possibly leading to prolonged irradiation and pronounced by-product formation.^{31,32} Pressurized mercury lamps are often used as light sources, although the emission of these high power lamps contains a considerable amount of UV light.³³ Expensive filter equipment is necessary for the selection of defined emission bands as well as an efficient heat management for avoiding thermal reaction pathways.

Those limitations can be overcome by the application of continuously running microreactors and modern light emitting diodes (LEDs) making this combination a versatile tool for photochemical applications.^{34,35} Microstructured equipment is able to contact reagents in gas and/or liquid streams more efficiently due to a strongly increased surface to volume ratio.^{36–39} The reaction volume inside a microreactor is very small and suited for an excellent process control. Hot spot

^a Fraunhofer ICT-IMM, Carl-Zeiss-Straße 18-20, 55129 Mainz, Germany.

E-mail: thomas.rehm@imm.fraunhofer.de

^b École Polytechnique Fédérale de Lausanne, EPFL-ISIC, Section 6, 1015 Lausanne, Switzerland

† Electronic supplementary information (ESI) available: Lab plant flow chart, sample taking device, HPLC analysis, UV/vis absorption spectra of the DHN and Juglone, oxygen gas–liquid mass transfer, TcPP synthesis, sensitized endoperoxide side-reaction, quantum efficiency calculation. See DOI: 10.1039/c6re00169f



formation becomes manageable by exact temperature control leading often to improved conversion and selectivity in the reaction.⁴⁰ Especially the irradiation of the reaction solution becomes easier. Thin film formation in microchannels or capillaries with a small inner diameter allows the full irradiation of the complete reaction volume inside the microchannels.⁴¹ The irradiation time can be exactly controlled by the flow rate of the reaction composition being pumped through the microstructures. These advantages offer many possibilities for improving in parallel the economic and the environmental sustainability of a chemical process.⁴² LEDs fundamentally enhanced the application range of photochemistry. Generating visible light of defined wavelength by low power consuming semiconductor materials has made LEDs to the light source of the future.^{43,44} LED technology necessitates neither filter equipment nor as intensive heat removal as it is the case for mercury lamps. LEDs are small-sized spot light emitters and are adaptable in arraying them according to the surface to be irradiated. These advantages make LEDs an ideal partner to continuously running microreactors.

Micro tubular-based reactors are an often used reactor type for continuous-flow photochemistry. Depending on the diameter of the tubular channel these reactors are used for process optimization on smallest scale (50–100 μL reactor volume) or for scale-up approaches under optimized conditions (up to hundreds of mL reactor volume). Takahashi *et al.* use a series of two small-sized glass reactors of 50 and 100 μL for the synthesis optimization of vitamin D₃ from the provitamin D₃ *via* a combined photochemical and thermal synthesis route.⁴⁵ Zeitler *et al.* performed a photocatalytic reductive dehalogenation in a glass microreactor of 100 μL reactor volume yielding a reaction time decrease from 12 h to 40 s compared to the batch synthesis.⁴⁶ As one example for a small scale-up Ryu *et al.* connected two 8 mL glass-covered stainless-steel microchannel reactor in series for the synthesis of a key intermediate for myriceric acid A. Using Black light irradiation this set-up produced the desired compound on the gram-scale.⁴⁷ Booker-Milburn *et al.* introduced a capillary photoreactor design in 2005 using a FEP capillary wrapped around an immersion well with a UV light source inside.^{48,49} Using capillaries with inner diameters of 3–4 mm allow large reactor volumes for scaling-up reactions without clogging. Photocycloadditions were transferred from batch to continuous-flow mode proving the superior productivity of the flow reactors on the multi-gram scale. This reactor concept was then used as model⁵⁰ for a large number of custom-built flow reactors for purely liquid or gas–liquid reactions like C–H functionalization of tertiary arylamines,⁵¹ addition of glycosyl radicals into acrolein,⁵² azide reduction and bromination of alcohols,⁵³ oxidative cyanation of amines,⁵⁴ the Stadler–Ziegler synthesis for aromatic C–S bond formation,⁵⁵ trifluoromethylation of heterocycles with gaseous ICF₃,⁵⁶ or photooxygenation during juglone synthesis.⁵⁷ Tubular-based reactors enable segmented flow of gas and liquid slugs inside the channels or capillaries allowing efficient gas mass trans-

fer into the liquid phase, while the reaction solution is fully illuminated. These conditions have been also used for the generation of singlet oxygen for the continuous-flow synthesis of artemisinin. Seeberger *et al.* modified and transferred the photooxygenation of dehydroartemisinic acid into flow mode.^{58,59} Recently, the process was enhanced by Rossen, Poliakov and George with the sensitizer immobilized on a ion exchanger resin.⁶⁰

Falling film reactors have been also applied to continuous-flow photochemistry. This type of reactors uses gravity to transport the reaction solution from the top of a glass barrel as thin film over the barrel's flat surface to its bottom. Griesbeck *et al.* combined this falling film reactor setup with a strong XeCl excimer lamp for irradiating the liquid film with UV light. A microstructured version of the falling film reactor was developed in order to decrease the liquid film thickness from millimetres to micrometres. The smaller thickness allows a strongly improved gas mass transfer into the liquid phase. Originally, the falling film microreactor (FFMR) was designed and successfully used as solvent evaporator and for highly exothermic reactions like hydrogenation of nitro groups.^{61–64} But its design (*vide infra*) also allows the selective irradiation of the reaction solution in its microstructure. Up to now the FFMR has been applied to three continuous-flow photochemical reactions: a) photochlorination of aromatic isocyanates,⁶⁵ b) cycloaddition of singlet oxygen into cyclopentadiene,⁶⁶ and c) cycloaddition of singlet oxygen into 1,5-dihydroxynaphthalene for the synthesis of juglone.⁶⁷

All reactions performed up to now lack of a detailed investigation of the process parameters of a continuous-flow photochemical reaction performed in the FFMR. To close this gap, we investigated the juglone synthesis as benchmark reaction performed in the FFMR with two sensitizers having different absorption wavelengths. The impact of the light source regarding the irradiation wavelength and the LED power are evaluated as well as the oxygen partial pressure and substrate concentration. Finally the architectural impact of the reactor will be evaluated as well. The quantum efficiency q , the productivity I_p and the space time yield STY are calculated for comparing the FFMR with other reactor types.

Results and discussion

Photomicroreactor and light sources

The FFMR is composed of a back plate carrying the heat exchanging unit and the inlet and outlet for the reaction solution (details can be found in Fig. S1 and S2 in the ESI†). It is followed by the reaction plate as the first key element of the reactor. On the front side of the reaction plate microchannels with different geometries for each plate have been incorporated by wet chemical etching (Table 1).

The reaction solution is fed through the top slit and pulled into the microchannels by gravity and capillary forces. Depending on the flow rate and viscosity of the solution thin films can be generated far below 100 μm . The reaction plate



Table 1 Geometrical data of FFMR reaction plate microchannels

Abbreviation	Channel number N	Width $w_{ch}/\mu\text{m}$	Depth $d_{ch}/\mu\text{m}$	Length l_{ch}/mm	Irradiated length $l_{irr\ ch}/\text{mm}$
FFMR_600	32	600	200	78	54
FFMR_1200	16	1200	400	78	54

is fixed with a mask and sealing gasket followed by the top plate of the reactor housing. This last part contains the second key element of the reactor, an inspection window made of borosilicate glass (visible to irradiation: length $l_{irr} = 54$ mm, width $w_{irr} = 29$ mm, thickness: 12 mm). Due to the high transparency of borosilicate glass for visible light (90% for wavelengths of $370\text{ nm} \leq \lambda \leq 2000\text{ nm}$)⁶⁸ it is most suitable for irradiating the thin film on the reaction plate with external light sources. Beside the inspection window the top plate carries the inlet and outlet devices of the gas phase, which is typically fed from bottom to top of the reactor. This arrangement allows a counter current flow of the gas stream against the liquid flow resulting in an excellent gas liquid contacting and mass transfer.

LEDs were installed as energy efficient sources for irradiation. Violet, royal blue, green and cold-white LEDs (GeTian Opto-Electronics CO., LTD.) were used in order to allow the selective irradiation of different photocatalysts (Table 2, Fig. 4).

An aluminium body was designed as heat sink carrying two LEDs (Fig. 1). These LED arrays can be fixed magnetically in front of the inspection window of the FFMR allowing a quick exchange of the array maintaining the same light path properties for all LEDs used in this work. Due to the LED's wide angle of radiation of 120° the complete reaction plate inside the reactor was irradiated. Special attention was drawn on a light-proof design of the LED array due to two reasons. First, the use of high-power irradiation, especially close to the UV region, requires safety precautions for protecting the eyesight of the experimenter. Second, light-tightness is the prerequisite for the full availability of all photons produced by the LED array inside the reactor to be used for the photochemical reaction.

Lab plant design

The lab plant was equipped with standard components for the complete set of experiments (Fig. 2). A water-filled thermostat (Julabo F10 and HC E07) was used to maintain the

Table 2 Spectral parameters of LED

LED	Emission maximum λ_{max}/nm	FWHM ^a /nm
Violet	410	22
Royal blue	455	24
Green	520	37
Cold-white (blue part)	442 (sharp)	22
Cold-white (red part)	550 (broad)	123

^a Full width at half maximum of the spectral band.

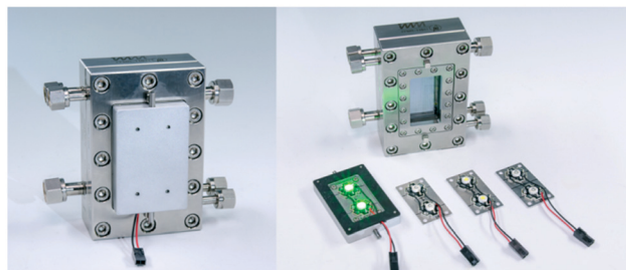


Fig. 1 Falling film microreactor with a magnetically attached heat sink carrying exchangeable LED arrays (©Fraunhofer ICT-IMM).

temperature of the FFMR at $20\text{ }^\circ\text{C}$. Synthetic air or pure oxygen (Linde AG) was released from a pressurized steel cylinder and broad to 1 bar application pressure with an appropriate pressure regulator (Linde AG, FMD 300-18). The fine regulation of current and voltage for the LED arrays were done with a standard power supply (Basetech BT-305). Two piston pumps (Chemyx Inc., Nexus 5000) were used to feed and withdraw the liquid phase from the FFMR. The gas flow rate was regulated by a mechanical rotameter (Baley, Fisher & Porter, G10A6142DA131X100). A flow chart can be found in Scheme S1 in the ESI.†

For sample taking a device was implemented between the reactor outlet and the withdrawing pump using two needles inside a HPLC vial. The feeding needle was placed at the bottom of the vial, whereas the withdrawing needle was placed 12 mm above the vial's bottom. This arrangement allows the continuous filling of the HPLC vial during the process with a defined volume of reaction solution. For each HPLC sample a



Fig. 2 Lab plant used for photochemical transformations with a falling film microreactor as core element.



new vial was plugged onto the stationary mounted needles (see Fig. S3 in the ESI†). Details on the analytical HPLC method can be found on page S5 of the ESI.†

Benchmark reaction and sensitizers

The photooxygenation of DHN has been performed in several types of flow reactors so far. Beside classical immersion well reactors equipped with mercury pressure lamps also sunlight collecting and concentrating reactors were used for the continuous-flow synthesis of juglone,^{69–71} whereas microscale reactors like capillary reactors overcome insufficient irradiation.^{57,72}

The key step in the photooxygenation of 1,5-dihydroxynaphthalene (DHN) to juglone is the *in situ* formation of singlet oxygen that pursues a [4 + 2] cycloaddition with DHN.^{73–75} The resulting endoperoxide is unstable and is instantaneously transformed to juglone and water (Fig. 3). Singlet oxygen can be produced under harsh chemical conditions⁷⁶ (being often not compatible with the stability of substrate molecules) or *via* a mild photocatalytic process.⁷⁷ The latter case needs a sensitizer like rose bengal (RB) or *meso*-tetraphenylporphyrine (TPP) exhibiting a triplet state energy greater than 95 kJ mol⁻¹.⁷⁸ After irradiation with light of appropriate wavelength the sensitizer is excited first to its singlet state which undergoes intersystem crossing (ISC) to the triplet state.⁷⁸ With a triplet oxygen molecule in close vicinity to the sensitizer energy transfer is possible yielding the highly reactive singlet oxygen species.⁷⁹ Although RB achieves a high quantum yield for singlet oxygen formation, it is known that this naturally occurring dye decomposes quite rapidly by photobleaching.⁸⁰ In the case of TPP the triplet quantum yield is slightly lower, but the photostability of the dye is considerably higher.⁸¹

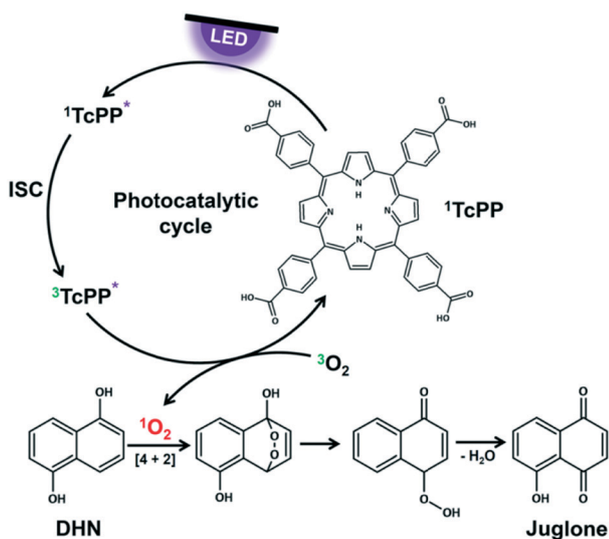


Fig. 3 Photocatalytic cycle of TcPP for singlet oxygen formation as key step in juglone synthesis.

To ensure comparability between the experiments both sensitizers are used under the same reaction conditions. RB is a salt which dissolves very good in aqueous solutions, but neither DHN nor TPP are soluble in water. In consequence, 2-propanol was chosen as environmental friendly solvent for all experiments.⁸² It has a cut-off wavelength of 240 nm ensuring that no interactions take place between the incident visible light and the reaction medium. The nonpolar TPP was substituted by its derivative *meso*-tetracarboxyphenylporphyrine (TcPP) containing four free carboxylic acids in *para*-position of the phenyl residues.⁸³ Despite a small bathochromic shift of the *Soret* band, these functional groups do not disturb the UV/vis absorption properties of the monomeric porphyrine core in polar solutions, but enhance the solubility of the sensitizer for its use in the chosen solvent.⁸⁴ Details on the Synthesis of TcPP can be found in the ESI† on page S6.

Definitions and control experiments

For a clear comparison of all experiments throughout this work a set of physical and chemical reaction parameters were defined (Table 3). Variations of these parameters due to a detailed investigation of their impact on the photocatalytic process are mentioned accordingly.

The main parameters varied in this work are the type of sensitizer (rose bengal or TcPP) and the light source (green, royal blue, violet, and cold-white). For an easier description of the reaction conditions the following sensitizer-light source combinations are abbreviated as defined in Table 4.

A control experiment has been performed in order to assess the necessity of a sensitizer for a successful realization

Table 3 Defined standard solution and standard reaction conditions for the continuous-flow synthesis of juglone in a FFMR

Parameter	Value
Substrate concentration c_{DHN}	$10 \times 10^{-3} \text{ mol L}^{-1}$
Sensitizer concentration c_{sens}	$0.5 \times 10^{-3} \text{ mol L}^{-1}$
Solvent	2-Propanol
Liquid flow rate f_l	0.16 mL min^{-1}
Gas flow rate f_g	100 mL min^{-1}
Reactor temperature T_{FFMR}	$20 \text{ }^\circ\text{C}$
Reactor pressure p_{FFMR}	Atmospheric pressure
LED array amperage I_{LED}	350 mA

Table 4 Abbreviations of sensitizer-light source combinations used in this work

Sensitizer	Light source	Abbreviation
Rose bengal	Cold-white	RB-cw
	Violet	RB-v
	Royal blue	RB-b
	Green	RB-g
<i>meso</i> -Tetracarboxy-phenylporphyrine	Cold-white	TcPP-cw
	Violet	TcPP-v
	Royal blue	TcPP-b
	Green	TcPP-g



of the juglone synthesis. A standard solution without sensitizer was used under standard conditions with a cold-white LED providing the broadest emission range. As expected no conversion to juglone was observed due to the absence of singlet oxygen solely produced by an appropriate sensitizer. Beside this prove any photochemical decomposition pathways of the starting material could be ruled out as seen by the HPLC analysis. Further control experiments were performed in order to assess the photochemical decomposition of juglone during the irradiation. Compared to DHN juglone absorbs light in the visible spectrum which might lead to consecutive reactions (for the UV/vis absorption spectra, see Fig. S6 in the ESI†). Commercially available juglone was used as starting material in the standard solution instead of DHN. Four experiments were performed with RB-cw, RB-g, TcPP-cw and TcPP-v under standard conditions. HPLC analysis confirmed that no decomposition of juglone or any follow-up reactions occurred in these experiments.

Influence of light source on sensitizer excitation

As first parameter the emission wavelength was evaluated for both sensitizers with pure oxygen on a FFMR_1200 plate under standard conditions. RB and TcPP have a specific UV/vis absorption pattern which result in unique emission-absorption overlaps for each LED array (Fig. 4). Each overlap is representative for the excitation of the sensitizers and in consequence for the singlet oxygen formation.

In the case of RB the gradual increase in conversion correlates with the increasing overlap of light emission and dye absorption. The best yields were obtained for the cold-white and the green LED (Fig. 5). The broad emission of the cold-white LED covers the whole absorption band of RB in the visible region between 470 nm and 600 nm. Its peak emission is in perfect overlap with the maximum absorption of RB at 550 nm. The green LED emission spectrum overlaps only in part with the RB absorption, particularly in the low absorbing region between 470 nm and 550 nm. The small overlap in RB-v and RB-b is clearly confirmed by the low conversion of 9% and 15%, respectively. Concerning the product selectivity, RB-g achieved the highest selectivity of $S = 93\%$, whereas RB-cw reached a slightly lower selectivity of $S = 87\%$. Although RB-cw results in higher conversion compared to RB-g it was

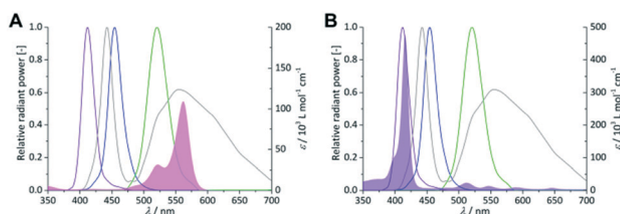


Fig. 4 Coloured lines indicate the spectral emission bands of the violet, royal blue, green and cold-white LEDs (left Y-axis = relative radiant power). UV/vis absorption spectra of $0.25 \times 10^{-3} \text{ mol L}^{-1}$ RB (A; pink area) and $1 \times 10^{-6} \text{ mol L}^{-1}$ TcPP (B; violet area) in 2-propanol (right Y-axis = molar extinction coefficient).

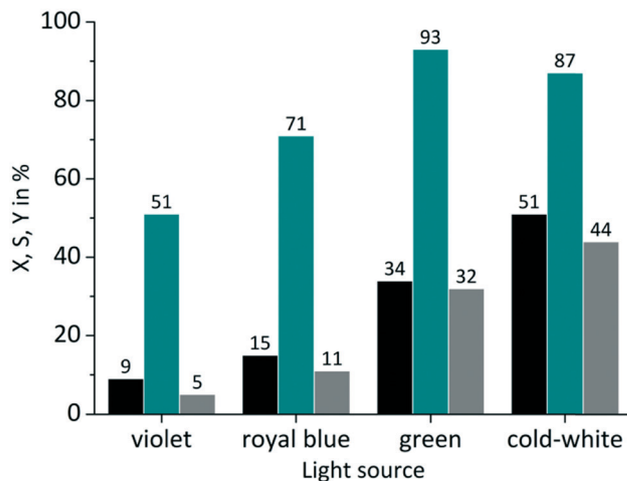


Fig. 5 Impact of emission wavelength on conversion and selectivity in juglone synthesis with RB and oxygen on a FFMR_1200 plate (■ conversion X; ■ selectivity S; ■ yield Y).

decided to the favour the higher selectivity with RB-g for further experiments.

The same experiments were performed with TcPP (Fig. 6), which exhibits a split visible light absorption pattern with the *Soret* band ($\epsilon_{\text{max}} = 416 \text{ nm}$) and the *Q* bands ($\epsilon_{\text{max}} = 511, 547, 589, 646 \text{ nm}$). As it was the case for RB-g and RB-cw the best spectral overlap gives the highest conversion. TcPP-v achieves a conversion of $X = 76\%$. The excellent overlap of the violet emission band with the strongly absorbing *Soret* band of the porphyrin core strongly contributes to high conversions for TcPP-v. Due to a shift to lower wavelengths and in consequence due to a less good spectral overlap TcPP-b only achieves a conversion of $X = 18\%$. In the case of TcPP-g the green light emission shows a very good overlap with the *Q* bands of the porphyrin core resulting in a conversion of $X = 19\%$. The emission of the cold-white LED allows a small overlap with the strongly absorbing *Soret* band and a very

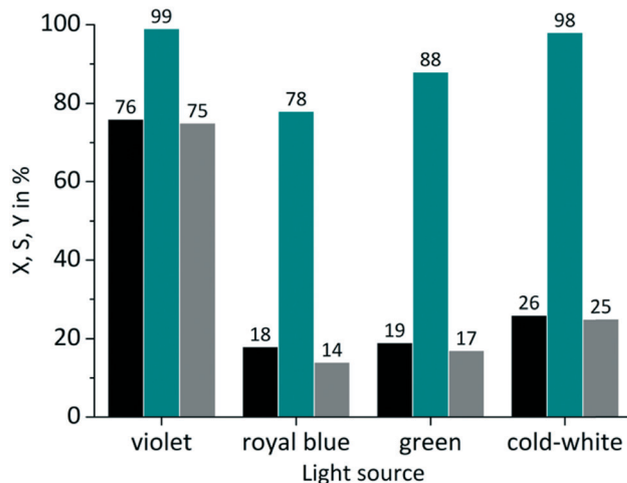


Fig. 6 Impact of emission wavelength on conversion and selectivity in juglone synthesis with TcPP and oxygen on a FFMR_1200 plate (■ conversion X; ■ selectivity S; ■ yield Y).



good overlap with the Q bands of the porphyrin core. This combination results for TcPP-cw in a conversion of $X = 26\%$. Interestingly TcPP allows achieving higher selectivity than RB reaching nearly 100% for TcPP-v and TcPP-cw, 88% for TcPP-g and 78% for TcPP-b.

Influence of LED power

In close conjunction to the first parameter the impact of the light power was evaluated in a second set of experiments. The amperage I_{LED} of the LED arrays was varied from 175 mA over 350 mA (standard value) to 530 mA. These values correspond to a power P_{el} of 1.1 W, 2.38 W and 3.6 W respectively for the green LED array used in the tests. RB was applied on a FFMR_1200 plate under standard conditions with oxygen gas (Fig. 7).

The variation of the power has an impact on both the conversion and the selectivity. The increase in conversion can be attributed to the larger number of photons available with increasing power for singlet oxygen formation. Selectivity changed less strongly with increasing LED power, but with the same trend to higher values. In the case of switching the electrical power from 1.1 W to 2.38 W a relative increase in conversion of 48% and selectivity of 6% is given. The increase of electrical power from 2.38 W to 3.6 W (approx. 51%) results in a relative increase of only 9% and 2% for conversion and selectivity. These rather small differences are an important result for the optimisation of the photochemical process regarding its energy consumption. The low increase in conversion points to the fact that the larger number of available photons at 3.6 W are not used for the photochemical process.

Influence of oxygen partial pressure

The availability of dissolved oxygen is an important key factor for the singlet oxygen formation during the juglone synthesis. The solubility of oxygen in 2-propanol is given as its molar

fraction with a value of $7.88 \times 10^{-4} \text{ mol mol}^{-1}$ at 20 °C and 1.013 bar.⁸⁵ Applying Henry's law results under these conditions in an oxygen concentration $^{O_2}c_{\text{oxygen}} = 10 \times 10^{-3} \text{ mol L}^{-1}$ (see Fig. S7 in the ESI†). The impact of the dissolved oxygen concentration was examined by the application of either pure oxygen ($^{O_2}p_{\text{oxygen}} = 1.013 \text{ bar}$) or synthetic air with a fivefold lower oxygen content ($^{\text{air}}c_{\text{oxygen}} = 2 \times 10^{-3} \text{ mol L}^{-1}$; $^{\text{air}}p_{\text{oxygen}} = 0.202 \text{ bar}$). RB was used as sensitizer on a FFMR_1200 plate under standard conditions. As can be seen in Fig. 8 the use of synthetic air has only a small negative impact on the conversion rate compared to pure oxygen. The slightly decreased conversion for RB-g by 2% and for RB-cw by 7% can be attributed to a lower oxygen concentration in the liquid phase due to the decreased oxygen partial pressure in the synthetic air gas phase. But in general the photochemical process does not seem to be mass transfer limited by the oxygen gas phase.

In the case of selectivity the use of synthetic air has a far more negative impact. Especially with RB-g the selectivity drops by 26%. In all other cases the selectivity decreases by a value of approx. 13%. One reason for this moderate selectivity might be a competing reaction that involves the sensitized conversion of the endoperoxide intermediate. Bicyclic endoperoxides are known to be cleaved by appropriate irradiation resulting in oxygen radicals.⁸⁶ These radicals can react intramolecularly with a double bond yielding epoxides.⁸⁷ The existence of a sensitizer can promote this reaction,⁸⁸ if the sensitizer is not quenched upfront by another reaction pathway like the singlet oxygen formation (see Scheme S2 in the ESI†). Due to the absence of excess oxygen in synthetic air RB might promote this competing reaction which is less pronounced in an oxygen-rich solution.

Influence of reaction plate architecture

The FFMR design comprises different reaction plate architectures (Table 1). As known from literature each channel

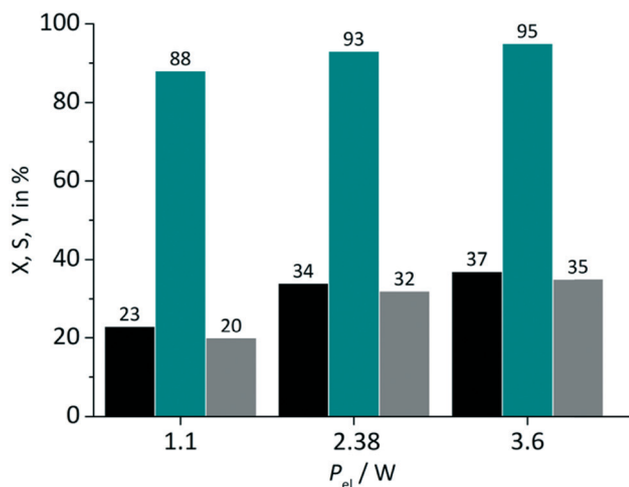


Fig. 7 Impact of light power on conversion and selectivity in juglone synthesis with RB and oxygen on a FFMR_1200 plate (■ conversion X; ■ selectivity S; ■ yield Y).

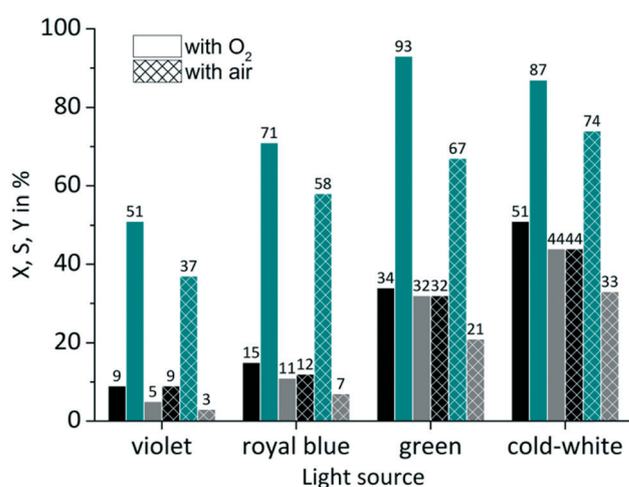


Fig. 8 Impact of oxygen partial pressure on conversion and selectivity in juglone synthesis with RB on a FFMR_1200 plate (■ conversion X; ■ selectivity S; ■ yield Y).



geometry can result in different thin film geometry.⁸⁹ Variations in thin film thickness have an impact on the gas-liquid phase contacting behaviour and in consequence on the conversion and selectivity of the examined reaction. The thin film formation can be investigated with a confocal laser microscope.^{38,90,91} With this method the experimental comparison of uncoated FFMR_600 with FFMR_1200 plates gave at the same flow rate a greater film thickness for the FFMR_600 than for the FFMR_1200 plate. A greater film thickness results in a longer residence time at the same flow rate applied for both plates offering the same channel area ($A_{\text{channels}} = 1497.6 \text{ mm}^2$). This point is reflected by the results obtained in this experiment. For TcPP-v and RB-g a relative increase in conversion of 15% and 25%, respectively, is obtained using the plate with the smaller channel architecture (FFMR_600) providing a greater film thickness and a longer residence time (Fig. 9). Selectivity increases as well, but not as consistent as for the conversion. These results indicate that a longer residence time is beneficial for the photochemical conversion.

Influence of substrate inlet concentration

Three DHN inlet concentrations were evaluated ($5, 10, 20 \times 10^{-3} \text{ mol L}^{-1}$) keeping the amount of sensitizer constant at $5 \times 10^{-4} \text{ mol L}^{-1}$. All experiments were performed with synthetic air as oxygen source. For both RB-g and TcPP-v the conversion decreases with increasing DHN concentration (Fig. 10). This trend can be attributed to the higher relative amount of sensitizer and oxygen at lower substrate concentration.

With increasing DHN concentration the values for the selectivity seem to run into a plateau with approx. $S = 70\%$ for RB-g and $S = 88\%$ for TcPP-v. The higher selectivity for the lowest DHN concentration may result also from the competing reaction pathways between singlet oxygen formation and

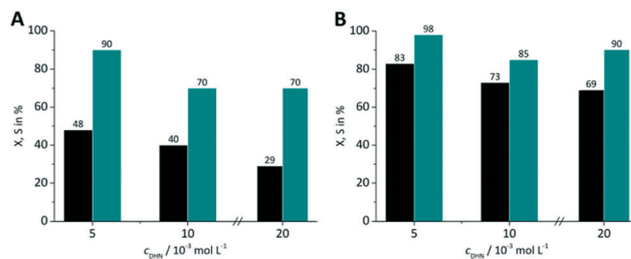


Fig. 10 Impact of DHN inlet concentration on conversion and selectivity in juglone synthesis with (A) RB-g and (B) TcPP-v on a FFMR_600 plate with air (■ conversion X; ■ selectivity S).

sensitized endoperoxide conversion. The more sensitizer and oxygen are available for singlet oxygen formation the less pronounced is the endoperoxide side reaction.

Optimization of reaction parameters

With the experimental results obtained so far the best combination was used for optimization. This was the case for TcPP-v with oxygen gas on a FFMR_600 plate under standard conditions ($X = 94\%$, $S = 99\%$). This arrangement was used for applying different LED power (Fig. 11A) and liquid flow rates (Fig. 11B). In the first case a lower LED power resulted in a significant relative loss in conversion of 25%, whereas with a higher LED power the selectivity decreased by 6% close to full conversion. Hence, the tests with varying flow rates were performed with the medium LED power corresponding to $P_{\text{el}} = 2.45 \text{ W}$. As expected, with the lowest flow rate of $f_l = 0.12 \text{ mL min}^{-1}$ the highest conversion was achieved yielding $X = 97\%$ with $S = 99\%$. With increasing flow rate the conversion decreased steadily to $X = 65\%$ at $f_l = 0.44 \text{ mL min}^{-1}$ maintaining an excellent selectivity of $S = 99\%$.

Theoretical part

In this section the correlation between the hydrodynamic behaviour of the reactor and the irradiation by the external LED arrays are presented and discussed for the experimental results with TcPP-v and RB-g. This section is split into two parts. In the first part the hydrodynamic and general parameters of the FFMR and the LED arrays are mathematically described and presented. The thickness of the liquid thin film

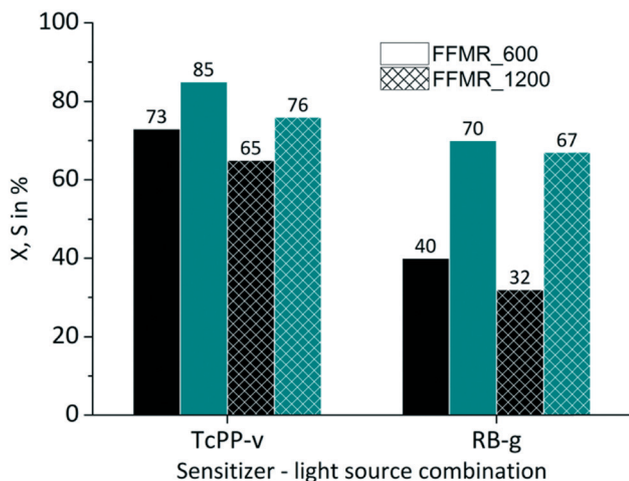


Fig. 9 Impact of plate architecture on conversion and selectivity in juglone synthesis with TcPP-v and RB-g with air (■ conversion X; ■ selectivity S).

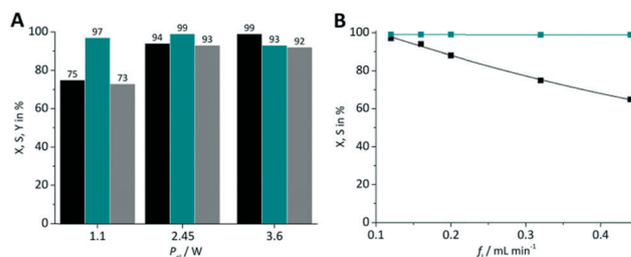


Fig. 11 Impact of (A) LED power and (B) flow rate on conversion and selectivity in juglone synthesis with TcPP-v on a FFMR_600 plate and oxygen (■ conversion X; ■ selectivity S, ■ yield Y).



inside the microchannels of the reaction plate is calculated as the main hydrodynamic parameter. The literature-known Nusselt eqn (1) allows a good approximation of the thin film thickness δ in the microchannels:^{92–94}

$$\delta = \sqrt[3]{\frac{3 \times f_l \times \mu_l}{N \times \rho_l \times w_{ch} \times g}} \quad [\mu\text{m}] \quad (1)$$

with f_l as the liquid flow rate (0.16 mL min⁻¹), μ_l as the dynamic viscosity of 2-propanol at 20 °C (2.3×10^{-3} kg m⁻¹ s⁻¹), ρ_l as the density of 2-propanol at 20 °C (0.786 g mL⁻¹) and g as the gravitational constant (0.981 m s⁻²). The parameters N and w_{ch} describe the number of microchannels and their width.⁹⁵ With eqn (2) the liquid phase volume V_{irr} inside the irradiated part of the microchannels can be calculated, the appropriate residence time τ with eqn (3):

$$V_{irr} = \delta \times l_{irr\ ch} \times w_{ch} \times N \quad [\text{mL}] \quad (2)$$

$$\tau = \frac{V_{irr}}{f_l} \quad [\text{s}] \quad (3)$$

Under standard reaction conditions the thickness of the liquid film is approx. 50 μm with a mean residence time of 19.4 s inside the irradiated microchannels. Based on the dimension of the inspection window of the FFMR the irradiated surface on the reaction plate inside the reactor can be calculated with eqn (4), whereas the irradiated surface of the microchannels can be calculated with eqn (5):⁹⁵

$$A_{irr} = l_{irr} \times w_{irr} \quad [\text{mm}^2] \quad (4)$$

$$A_{irr\ ch} = N \times l_{irr\ ch} \times w \quad [\text{mm}^2] \quad (5)$$

The irradiated microchannel area is with 1037 mm² approx. 65% of the entire irradiated area since the embossments between the microchannels are not wetted with the liquid phase. Table 5 summarizes the results of the calculation for the hydrodynamic parameters for the FFMR including the general parameters for the irradiated surfaces inside the microreactor.

In the case of the LED arrays the radiant power P_{LED} of the array is either given in the technical data sheet or can be calculated with this data.^{96,97} The luminescent efficiency η_{lum} of the LEDs can be calculated with eqn (6):

$$\eta_{lum} = \frac{P_{LED}}{P_{el}} \quad (6)$$

Table 5 Hydrodynamic and general parameters for the FFMR

Thin film thickness δ	50 μm
Irradiated volume V_{irr}	0.052 mL
Residence time τ	19.4 s
Irradiated area in FFMR A_{irr}	1599 mm ²
Irradiated area in microchannels $A_{irr\ ch}$	1037 mm ²

with P_{el} as the electrical power of the LED array at a current I_{LED} of 350 mA. With a transmission $T = 0.9$ for the borosilicate glass of the inspection window the radiant power available inside the FFMR can be calculated as well (eqn (7)):⁶⁸

$$P_{av\ LED} = 0.9 \times P_{LED} \quad [\text{W}] \quad (7)$$

Table 6 summarizes the results of the calculation for the general parameters for the LED arrays used with the FFMR.

In the second part the photophysical parameters of the incident light and its interaction with the photocatalyst are mathematically described and presented (details and numbers of the calculation can be found on page S10 and S11 of the ESI†).^{31,98} It has to be noted that the following equations and calculations reflect an idealized process in which the incident light trespasses the thin film just once. Any scattering effects or reflections from the reactor chamber walls resulting in multiple light transits through the solution are not considered.⁹⁹ The relative radiant power $P_{rel\ \lambda}$ of the LED arrays was extracted for each wavelength from the technical data sheet of the LED manufacturer. The decadic molar extinction coefficient was obtained from the UV/vis absorption spectra of both dyes. The density function of the light source g_λ given by eqn (8) is defined as normalization factor for the calculation of the entire radiant power φ_λ per wavelength in eqn (9):

$$g_\lambda = \frac{P_{av\ LED}}{\sum_{\lambda\ min}^{\lambda\ max} P_{rel\ \lambda}} \quad [\text{W}] \quad (8)$$

$$\varphi_\lambda = P_{rel\ \lambda} \times g_\lambda \quad [\text{W nm}^{-1}] \quad (9)$$

The available radiant power per wavelength in the microchannels $\varphi_{\lambda\ ch}$ is defined by the ratio of the entirely irradiated surface to the irradiated surface of the microchannels (eqn (10)):

$$\varphi_{\lambda\ ch} = \varphi_\lambda \times \frac{A_{irr\ ch}}{A_{irr}} \quad [\text{W nm}^{-1}] \quad (10)$$

Based on the experimentally obtained decadic molar extinction coefficient ε_λ the Napierian extinction coefficient μ_λ can be calculated for each wavelength (eqn (11)):

$$\mu_\lambda = \varepsilon_\lambda \times c_{sens} \times \ln(10) \quad [\text{m}^{-1}] \quad (11)$$

Table 6 Available radiant power and luminescent efficiency of each LED array

	Violet LED	Green LED
Voltage U_{LED} at 350 mA	7 V	6.8 V
Electrical power P_{el}	2.45 W	2.38 W
Radiant power P_{LED}	0.84 W	0.37 W
Luminescent efficiency η_{lum}	34.3%	15.6%
Available radiant power $P_{av\ LED}$	0.76 W	0.34 W



Table 7 Calculated results for the absorption of available power and photons under standard reaction conditions

	TcPP-v	RB-g
Available power φ_{ch}	0.49 W	0.217 W
Absorbed power φ_{abs}	0.319 W	0.034 W
Energy efficiency η_E	0.13	0.014
Mean photon absorption efficiency $\bar{\eta}_{\text{photon abs}}$	0.29	0.19
Molar amount of available photons $n_{\text{photons ch } \tau}$	3.3×10^{-5} mol	1.8×10^{-5} mol
Molar amount of absorbed photons $n_{\text{abs photons } \tau}$	2.1×10^{-5} mol	2.9×10^{-6} mol

Eqn (10) and (11) can be used on base of the Lambert-Beer law for the calculation of the radiant power per wavelength absorbed by the sensitizer in the liquid thin film $\varphi_{\text{abs } \lambda}$ (eqn 12):

$$\varphi_{\text{abs } \lambda} = \varphi_{\lambda \text{ ch}} \times \left(1 - e^{(-\mu_{\lambda} \times \delta)}\right) \quad [\text{W nm}^{-1}] \quad (12)$$

With eqn (13) the photon absorption efficiency per wavelength $\eta_{\text{photon abs } \lambda}$ can be expressed by the ratio between the absorbed radiant power and the available radiant power per wavelength on the structured surface:

$$\eta_{\text{photon abs } \lambda} = \frac{\varphi_{\text{abs } \lambda}}{\varphi_{\lambda \text{ ch}}} \quad (13)$$

The results from eqn (10) and (12) can now be converted into the molar amount of available photons (eqn (14)) and molar amount of absorbed photons absorbed by the sensitizer per wavelength during residence time τ (eqn (15)):

$$n_{\text{photons ch } \lambda \tau} = \frac{\varphi_{\lambda \text{ ch}} \times \lambda \times \tau}{h \times c \times N_A} \quad [\text{mol nm}^{-1}] \quad (14)$$

$$n_{\text{abs photons } \lambda \tau} = \frac{\varphi_{\text{abs } \lambda} \times \lambda \times \tau}{h \times c \times N_A} \quad [\text{mol nm}^{-1}] \quad (15)$$

with λ as the specific wavelength, h as the Planck constant, c for the speed of light and N_A as the Avogadro constant. The results of eqn (10), (12), (14) and (15) represent the available and absorbed power and the molar amount of available and absorbed photons, respectively, for each wavelength. With eqn (16)–(19) these values are summarized in order to address the whole spectral emission and absorption range of the appropriate light source and sensitizer combination:

Table 8 Quantum efficiency q , productivity L_p and STY for the processes with TcPP-v and RB-g under standard reaction conditions

Entry		TcPP-v	RB-g
1	Quantum eff. q (air)	0.0095	0.008
2	Quantum eff. q (O ₂)	0.014	—
3	Productivity L_p (air)	0.06 mmol h ⁻¹	0.03 mmol h ⁻¹
4	Productivity L_p (O ₂)	0.09 mmol h ⁻¹	—
5	STY (air)	0.32 mol s ⁻¹ m ⁻³	0.14 mol s ⁻¹ m ⁻³
6	STY (O ₂)	0.48 mol s ⁻¹ m ⁻³	—

$$\varphi_{\text{ch}} = \sum_{\lambda_{\text{min}}}^{\lambda_{\text{max}}} \varphi_{\lambda \text{ ch}} \quad [\text{W}] \quad (16)$$

$$\varphi_{\text{abs}} = \sum_{\lambda_{\text{min}}}^{\lambda_{\text{max}}} \varphi_{\text{abs } \lambda} \quad [\text{W}] \quad (17)$$

$$n_{\text{photons ch } \tau} = \sum_{\lambda_{\text{min}}}^{\lambda_{\text{max}}} n_{\text{photons ch } \lambda \tau} \quad [\text{mol}] \quad (18)$$

$$n_{\text{abs photons } \tau} = \sum_{\lambda_{\text{min}}}^{\lambda_{\text{max}}} n_{\text{abs photons } \lambda \tau} \quad [\text{mol}] \quad (19)$$

The energy efficiency η_E can be described as the ratio between the absorbed power φ_{abs} against the electrical power P_{el} (eqn (20)):

$$\eta_E = \frac{\varphi_{\text{abs}}}{P_{\text{el}}} \quad (20)$$

Averaging the photon absorption efficiency over all wavelengths gives the mean photon absorption efficiency for the whole process (eqn (21)):

$$\bar{\eta}_{\text{photon abs}} = \frac{\sum_{\lambda_{\text{min}}}^{\lambda_{\text{max}}} \eta_{\text{abs } \lambda}}{\lambda_{\text{max}} - \lambda_{\text{min}}} \quad (21)$$

The calculated result shows that TcPP-v is more efficient in the energy conversion as well as in the photon absorption with $\eta_E = 0.13$ and $\bar{\eta}_{\text{photon abs}} = 0.29$ compared to $\eta_E = 0.014$ and $\bar{\eta}_{\text{photon abs}} = 0.19$ for RB-g. The stronger efficiency of TcPP-v can be attributed to the better overlap of the main emission band of the LED array and the absorption band of the porphyrine sensitizer. One has also to keep in mind that both sensitizers have been used with the same molar concentration of 5×10^{-4} mol L⁻¹ whereas the sensitizers differ in their extinction coefficients at their particular maximum absorption wavelength by a factor of approx. 4.7 ($^{416}\epsilon_{\text{TcPP}} = 476846 \text{ M}^{-1} \text{ cm}^{-1}$ vs. $^{562}\epsilon_{\text{RB}} = 109410 \text{ M}^{-1} \text{ cm}^{-1}$). Table 7



Table 9 Comparison of productivity L_p and STY for various combinations of sensitizers, white light sources and reactor types with $c_{\text{DHN}} = 10$ mM, RB with air and TcPP/TPP with oxygen gas

Entry		This work FFMR_1200	Lit. [67] FFMR_1200	Lit. [57] capillary reactor
1	Sens. LED	L_p [mmol h ⁻¹] 0.03	RB ^a 0.0015	—
2		STY [mol s ⁻¹ m ⁻³] 0.17	0.008	
3	CFL ^c	L_p [mmol h ⁻¹] —	0.006	—
4		STY [mol s ⁻¹ m ⁻³] —	0.04	
5	Sens. LED	TcPP ^a L_p [mmol h ⁻¹] 0.02	—	TPP ^b 0.74
6		STY [mol s ⁻¹ m ⁻³] 0.13		0.02

^a RB & TcPP concentration was 0.5 mM. ^b TPP concentration was 0.03 mM. ^c CFL = compact fluorescence lamp.

summarizes the results of the theoretical evaluation for TcPP-v and RB-g.

The quantum efficiency q , the productivity L_p and the space time yield STY represent three important parameters for comparing a (photochemical-) process, which has been performed in different reactor types. The quantum efficiency q can be calculated with eqn (22) *via* the ratio of the molar amount of juglone produced during the residence time against the molar amount of photons absorbed by the sensitizer:

$$q = \frac{f_1 \times c_{\text{DHN}} \times X \times S \times \tau}{n_{\text{photons ch } \tau}} \quad (22)$$

The productivity L_p represents the molar quantity of juglone produced per unit of time (eqn (23)):

$$L_p = f_1 \times c_{\text{DHN}} \times Y \quad [\text{mmol h}^{-1}] \quad (23)$$

The space time yield STY normalizes the productivity of the process to the liquid volume in the reactor (eqn (24)):

$$\text{STY} = \frac{L_p}{V_{\text{irr}}} \quad [\text{mmol h}^{-1} \text{ m}^{-3}] \quad (24)$$

All three parameters have been calculated for TcPP-v and RB-g used under standard reaction conditions with air on a FFMR_600 reaction plate and for TcPP-v also with oxygen gas (Table 8). In corollary to the efficiency results discussed above TcPP-v shows higher quantum efficiency than RB-g (Table 8, entry 1). Especially with oxygen gas TcPP-v shows an approx. 1.8 times higher quantum efficiency than RB-g. The same trend can be seen for the productivity L_p and the STY: RB-g/air < TcPP-v/air < TcPP-v/oxygen (Table 8, entry 3–6).

These results can be compared with literature data from Oelgemöller *et al.* for the FFMR⁶⁷ and from de Oliveira *et al.* for a capillary reactor⁵⁷ using eqn (23) and (24) (Table 9). For better comparability only results were used which were obtained with white light sources. In the case of the FFMR processes it can be clearly seen that L_p and STY have been increased by a factor of up to 21 solely due to an improved light

source architecture in our system (Table 9, entry 1–4). Furthermore, the capillary reactor can be seen as scale-up approach to the FFMR. A 37-fold increase in productivity (Table 9, entry 5) shows the strong impact of the different reactor architecture with larger reactor volume (25 mL) and higher liquid flow rate (0.75 mL min⁻¹).¹⁰⁰ In reverse, the STY of the FFMR is approx. 7 times higher than for the capillary reactor. Comparing the productivity of the capillary reactor with our optimized process in the FFMR using TcPP-v (Table 8, entry 4) instead of TcPP-cw (Table 9, entry 5) diminishes the factor of 37 to 8 for the productivity increase. This clearly indicates that an even higher productivity would be possible for the capillary reactor, if its light source would be optimized to the sensitizer, *i.e.* by using violet light for covering the *Soret* band of the porphyrin.

Conclusions

With the results of this work we present a detailed investigation of the utilization of a microstructured falling film reactor for a photochemical application with visible light. The excellent capability of the FFMR to contact gas and liquid phases is combined with high-power LED arrays for the photonic contacting of the liquid phase that contains the metal-free sensitizers. With this setup singlet oxygen is produced *in situ* for the photochemically catalysed oxygenation of 1,5-dihydroxynaphthalene to juglone. Detailed investigations were performed according the light source, radiant power, oxygen partial pressure, reaction plate architecture, and substrate concentration. After optimization of the reaction conditions a conversion rate as high as 97% at 99% selectivity was achieved for the porphyrine sensitizer irradiated with violet light under pure oxygen gas atmosphere. The photochemical process in the FFMR was also theoretically investigated. It could be shown that the quantum efficiency can be improved by chemical reaction optimization, *e.g.* switch to pure oxygen gas, and physicochemical optimization, with *e.g.* a good spectral overlap of the emission and absorption bands of the chosen photocatalyst and LED array. Productivity and space time yield were compared with literature data showing that both parameters can be increased dramatically with



specific adaptation of the light source to the reactor. Based on these results ongoing research activities investigate the variance in selectivity depending on the sensitizer-light source combination. Scale-up approaches and a more detailed observation of the irradiation characteristics inside the reactor chamber will be performed for a deeper understanding of the impact of scattering and reflection effects on the photochemical process in continuous-flow. This might allow in future an ecologically optimized combination of thin film thickness and full absorption of incident light at the main absorption wavelength of the sensitizer used.

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