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Efficient Singlet Oxygen Photogeneration by Zinc Porphyrin-Dimers Upon One- and Two-Photon Excitation.

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ABSTRACT

The development of photodynamic therapy at depth requires photosensitizers which have both sufficient quantum yield for singlet oxygen generation and strong two-photon absorption. Here we show that this can be achieved by conjugated linkage of zinc porphyrins to make dimers. We determined the quantum yield of generation of ${}^{1}O_{2}$, ϕ_{A} , by measuring emission at 1270 nm using a near infra-red streak camera and found it to increase from 15% for a single porphyrin unit to 27-47% for the dimers with a conjugated linker. Then, we measured the spectra of two-photon absorption cross section, σ_2 , by a focus-tunable Z-scan method, which allows for nondestructive investigation of lightsensitive materials. We observed a strong enhancement of the two-photon absorption coefficient in the dimers, especially those with an alkyne linker. These results lead to an excellent figure of merit for two-photon production of singlet oxygen (expressed by the product $\sigma_2 \times \phi_A$ in the porphyrin dimers, of around 3700 GM, which is very promising for applications involving treatment of deep tumors by photodynamic therapy.

1. INTRODUCTION

Photodynamic therapy (PDT) is a widely used cancer treatment, particularly for the treatment of superficial cancers such as many skin cancers1. PDT involves the use of light in combination with а photosensitizer to cause a photochemical reaction, leading to the generation of reactive oxygen species (ROS)2, which then kill neighboring tumor cells. Whilst most often used to kill cancer cells, the reactive oxygen species can also be used kill bacteria thereby providing a powerful alternative to antibiotics in the management of topical infections^{3, 4}. For example, in addition to cancers, PDT has been used in the treatment of acne⁵ (by

killing Propionibacterium acnes), periodontology⁶, treatment of viral lesions⁷ and age-related macular degeneration8. In order to efficiently photogenerate singlet oxygen, a material should fulfill a few criteria. It needs to absorb excitation light and have a triplet energy level higher than 397 kJ/mol to allow for efficient energy transfer to ground state oxygen and hence the generation of singlet oxygen. Moreover, it should have high triplet quantum yield and long triplet state lifetime². Despite these requirements, there are many materials capable of photogenerating singlet oxygen, they can be grouped into: (1) the organic dyes and

aromatics⁹; (2) the porphyrins, chlorins, phthalocyanines, and related macrocycles¹⁰; (3) semiconductors¹¹; and (4) metal nanoparticles^{12, 13}. Among these photosensitizers, porphyrin derivatives attracted significant attention and some of them are already in clinical use.

Photodynamic therapy has been widely used to kill tumor cells, which are located close to skin surface, due to the fact that the photosensitizers are excited with wavelengths which are strongly absorbed and scattered by the tissue. PDT could treat a much wider range of conditions if it could work at greater depth which could be achieved by using excitation

wavelengths biological the in transparency windows of tissue at 650-950 nm and 1000-1250 nm¹⁴. As only few photosensitizers have linear absorption band in this wavelength region¹⁵, two-photon excitation in which longer wavelength photons are two

simultaneously absorbed is a natural way to achieve the desired excitation energy with longer wavelength light that can further The penetrate into tissue. photosensitizers which aim to be used in two-photon excited PDT should be characterized by high quantum yield of singlet oxygen generation, ϕ_{Δ} , and large two-photon absorption cross-sections.

1

Typical porphyrin derivatives¹⁶⁻¹⁸, even those accepted for clinical use like Photofrin[®], Verteporfin[®] or Foscan[®] are indeed efficient singlet oxygen generators but suffer from low two-photon absorption cross-sections, on the order of 1 GM^{19, 20}. In that context, there is a need to design and synthesize new molecules with excellent figure of merit for two-photon production of singlet oxygen (expressed by the product $\sigma_2 \times \phi_A$). It is also relevant to

consider this figure of merit when normalized by molecular weight i.e. $\sigma_2 \times \phi_A / M$, marked as $(\sigma_2 \times \phi_A)^*$ to facilitate comparison of large and small candidate photosensitizers. ln the case of porphyrins, in order to maximize the parameter $\sigma_2 \times \phi_\Delta$, insertion of a metal ion leads to a trade-off as it causes reduction of the singlet oxygen quantum yield²¹ and increase of the 2PA cross-section.

In the field of nonlinear optics, the general rules for molecular design to maximize 2PA cross-section exist in the literature and are supported by both theoretical and experimental work. It has

been shown that centrosymmetric quadrupolar compounds have larger dipolar values of than σ_2 their counterparts²². In addition, introducing strong electron donating (D) or electron withdrawing (A) groups to yield the structures like D-A-D or A-D-A further improves 2PA properties²³. Finally, the process of simultaneous absorption of two photons requires strong overlap of the π-orbitals over a large system, making it sensitive to molecular conformation changes, which may alter the electronic coupling.

The general structure-property rules can be applied to porphyrin based compounds. It has been shown that 2PA

properties may be tuned by metal insertion, adding electron donating or withdrawing groups and expanding the π -conjugated system by creating dimers

or higher order oligomers^{17, 23, 24}. In the present work we use metal containing porphyrins and explore the effect of increasing electron delocalization by conjugated links between macrocycles. In particular, we explore the photogeneration of singlet oxygen and two-photon absorption in a series of zinc porphyrins, either being a monomer (compound 1), or

a dimer, linked by ethyne (2) or di-ethyne link (3) or by a single bond (4), as depicted in Scheme 1.

There are numerous papers investigating either the two-photon absorption or singlet oxygen photogeneration properties of porphyrin

the combination of these two effects is presented²⁵⁻²⁷. Our goal was to carry out interdisciplinary research covering the nonlinear optical properties and singlet oxygen photogeneration, thus providing strong support to the results and conclusions concerning the structure-property optimization. Singlet oxygen

streak camera. The two-photon absorption properties were investigated by the focustunable Z-scan method, which has the important advantage over alternative methods of being quick, thereby avoiding photodegradation which is a particular consideration in the measurement of materials intended to cause photochemistry.

generation was studied using an infra-red

Scheme 1. Structures of investigated compounds.

2. EXPERIMENTAL SECTION

The molecules used in this work were synthesized and purified according to the protocols described in 28. The molecules were dissolved in chloroform (Sigma-Aldrich) and quartz cuvettes were filled with the solutions for the photophysical measurements. UV-vis absorption spectra were measured using a Perkin Elmer Lambda 1050 UV/Vis/NIR

spectrophotometer. The concentration of the solution was adjusted to reach optical densities suitable for the PL and emission from singlet oxygen experiments. Steadystate PL spectra were recorded using an automated spectrofluorometer (Fluorolog, Horiba Jobin-Yvon), with a 450 W Xenon lamp excitation source. The solution singlet oxygen quantum yield (SOQY) (ϕ_{Δ})

determined values using were tetraphenylporphyrin (H₂TPP) from Fisher Scientific in chloroform solution as a standard reference because of its wellknown SOQY of 55%²⁹. The direct emission of singlet oxygen at 1270 nm was detected by a NIR streak camera from Hamamatsu Photonics. The camera was working in Photon Counting mode and the experiment was automatically stopped after 50 000 frames for each sample. In order to prevent the degradation, the samples solutions were placed in a rotating cell, placed in such a way that the exciting beam was shifted from the center.

PHAROS regenerative amplifier (output wavelength 1028 nm, pulse duration 290 fs, repetition rate 5 kHz) was directed to an optical parametric amplifier (OPA, a Light **OPA** Conversion OPHEUS-F). The generated 828 nm, which was further frequency doubled to obtain 414 nm used to excite the samples. The output pulse duration was less than 100 fs. The quantum yield singlet of oxygen generation may be determined using the modified method used by Tanielian and Heinrich³⁰ and using equation 1²¹:

$$\phi_{\Delta x} = \phi_{\Delta r} \frac{A_r(\lambda_r)}{A_x(\lambda_x)} \frac{I(\lambda_r)}{I(\lambda_x)} \frac{D_x}{D_r} \frac{\tau_r}{\tau_s}$$
(1)

The beam from a Light Conversion

where ϕ_A stands for the quantum yield of singlet oxygen generation, the subscripts x and r refer to the compound to be characterized and reference to the solutions, respectively. A is the excitation wavelength, $A(\lambda)$ is the absorbance, $I(\lambda)$ is the relative intensity of the excitation light, *r* is the lifetime of singlet oxygen emission in the sample and reference solvent, and D is the integrated area under the emission spectra.

As, the samples and reference solutions are excited at the same conditions and they are dissolved in chloroform, the above equation can be simplified to:

$$\phi_{\Delta x} = \phi_{\Delta r} \frac{A_r(\lambda_r)}{A_x(\lambda_x)} \frac{D_x}{D_r}$$
 (2)

The two-photon absorption cross-sections were determined by f-scan method described in³¹. The laser system used for this study was a Quantronix Integra Ti:Sapphire regenerative amplifier (output wavelength: pulse duration: 130 fs; repetition rate: 1 kHz) that pumped an optical parametric amplifier (OPA, a Quantronix Palitra) to provide tunable excitation. The sample solution dissolving prepared by the was corresponding chromophore in chloroform at concentration of 0.5% (w/w) and placed in a glass cell with 1 mm optical pathlength. The measurements were carried out in a relative manner, calibrating all the data against f-scans carried out on a fused silica plate and taking into account the nonlinear signals obtained from a cell containing pure solvent.

3. RESULTS AND DISCUSSION

Steady-state electronic absorption.

The absorption and photoluminescence spectra of the investigated compounds were measured in dilute CHCl₃ solutions with 1×10⁻⁶ mol/L concentration.

Maximum absorption wavelengths and maximum emission wavelengths are listed in Table 1. The absorption spectra are displayed in Figure 1. For the monomeric

absorption bands are clearly visible. The Soret band is located at 414 nm and so called Q-band at 543 nm. For the dimers, the Q-bands are intensified they show bathochromic shift compared to molecule 1. Moreover, both Soret and Q-bands are much broader with additional peaks visible in the absorption spectra for compounds **2-4**. It is worth to mention that the stronger Q-band absorption is responsible for enhanced 2PA properties in the NIR. Fluorescence occurs from the Q-band, whichever excited. The band is fluorescence bands of compounds 2 and 3 show a bathochromic shift with respect

porphyrin, compound 1, (Figure 1a), two

as $B_x(0-0)$ transition. There are also other compound 1 (Figure 1a-c). The bathochromic shift of the absorption and B_x and B_y transitions which span a wide fluorescence bands can be explained by region to the blue from the $B_x(0-0)$ the decreased HOMO-LUMO gap due to transition. In the case of compound 4 the π-delocalization between the neighboring molecule is no longer planar due to steric Polarized absorption porphyrin units²⁴. hindrance, and so the dimerization does spectroscopy revealed that for the dimers not lead to increased electron linked with ethyne and di-ethyne bonds, delocalization. There is an almost equally transitions intense split in the Soret absorption and Qthe lowest energy were polarized along the molecular axis (x) and band emission of 4 (Figure 1d), which can can be called $Q_x(0-0)^{17}$. They are followed be explained by the excitonic coupling by vibronic satellites. In the Soret region, between the two parallel strong dipole there is a group of x- and y-polarized transitions of each porphyrin ring^{28, 32}. The transitions. Among them, magnitude of the absorption band split is the lowest equal to 2002 cm⁻¹ and the emission band energy and relatively narrow transitions is parallel to S₁←S₀, which can be assigned split is equal to 854 cm⁻¹.

The crystallographic structure analysis performed by another group yielded the torsional angle of the porphyrin dimer linked by a single bond to be 69.1(2)°33.

The torsional angle of the dimers linked by ethyne and di-ethyne bonds were

determined by DFT calculations to be zero, which means that the molecules are planar³⁴, which enhances both 2PA and singlet oxygen photogeneration due to faster intersystem crossing (S_1-T_1) , compared to twisted conformers³⁵.

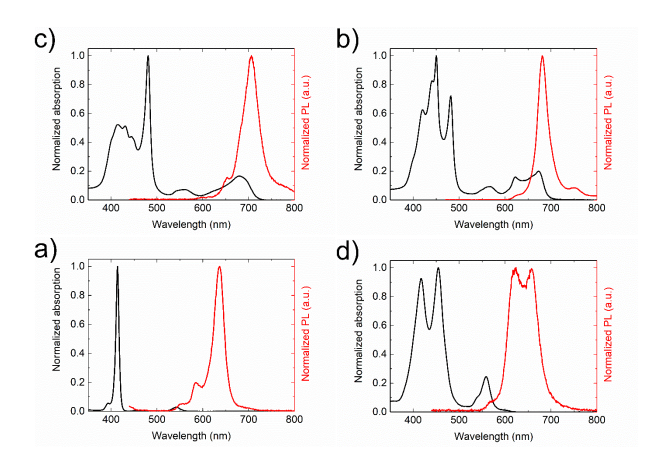


Figure 1. Normalized absorption (black curves) and steady-state fluorescence spectra (red curves) of investigated porphyrins. Figures a) to d) correspond to molecules 1 to 4, respectively.

Singlet oxygen detection.

The singlet oxygen quantum yield, ϕ_{A} , emission. The spectra obtained for sample 4 and the reference are shown in Figure 2. determined by recording direct was emission from singlet oxygen centered at Finally, the spectra were integrated to 1270 nm by a NIR streak camera. The obtain the D factors necessary to calculate transition from first excited singlet state the quantum yield of singlet oxygen ${}^{1}\Delta_{g}$ to ground state being triplet, ${}^{3}\Sigma_{g}^{-}$, is photogeneration. The values of ϕ_{Δ} of all spin forbidden². This is the reason for the investigated compounds are summarized in Table 1. The least efficient is the emission lifetime on the order of us, which strongly depends on the solvent²¹. monomeric unit (1). The dimerization

The signals were integrated from 8 to leads to enhanced properties of singlet $50 \, \mu s$ to obtain the spectra of 1O_2 oxygen photogeneration, regardless the

type of the link. One can notice that the value of ϕ_{Δ} for compound **4**, which is not conjugated due to steric hindrance, is approximately doubled, compared to 1. On the other hand, compounds 2 and 3, which have large π-delocalization and are close to planar, are characterized with much larger ϕ_{Δ} being equal to 35 and 47 %. findings These are good in agreement with other works on porphyrins dimers, e.g. with porphyrin units linked by butadiyne bond linked to TIPS-protected ethynyl groups³⁶. The determined ϕ_{Δ} of porphyrin dimer in DCM such was reported to be 37 %, which means that the structural design of compounds 2 and 3

are more efficient in terms of singlet oxygen production.

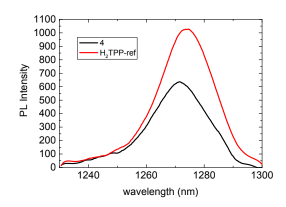


Figure 2. Singlet oxygen emission spectra of 4 and H_2 TPP, obtained by integrating the streak camera images from 8 to 50 μ s.

Table 1. Steady-state photophysical properties of the investigated compounds (1-4) and their singlet oxygen quantum yields. The positions of the most intensive bands are given in bold.

$\begin{array}{ c c c c c c c c c c c c c c c c c c c$	λ_{em} (nm)	$\phi_{arDelta} \ (\%)$	
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sample

	Soret band	Q- band		
1	393 414	542	554 585 637 700	15
2	420 441 450 482	566 622 673	627 681 751	35

	414		599	
3	431	562	654	47
	444	681	706	47
	481		785	
4	417	538	567	
	455	558	624	27
	433	600	658	

changes

the

measures

Two-photon absorption cross-section spectra.

transmission as a function of the sample position in Z direction and fit the data recorded on so called Open Aperture curve as described in the literature^{38, 41} to determine two-photon the value of absorption cross-section at the specific wavelength. In order to obtain reliable data by these two aforementioned techniques, the samples perfectly have photostable.

The spectra of two-photon absorption cross-section may be obtained by many methods, two-photon excited fluorescence (TPEF)³⁷ and fs Z-scan³⁸⁻⁴¹ being the most common nowadays. They strongly differ from each other, as in TPEF one detects fluorescence spectrum at various wavelengths and compares the integrated PL with a standard reference

with known PLQY and σ_2 . In Z-scan one

Upon photoexcitation with one- and two-photons the singlet oxygen which is generated by the investigated porphyrins causes their degradation. It is important to photosensitizer mention that each molecule can typically produce 10³-10⁵ molecules of ¹O₂ before being degraded through photobleaching by ¹O₂ or by some other process². Thus, it is crucial to minimize the time of irradiation especially with high power laser used to study the nonlinear effects. Hence we use the socalled "f-scan" technique which is a modification of the standard Z-scan method, to determine the spectra of two-photon absorption cross-section. In

the f-scan technique, instead of translating the sample, the focal point is shifted by an electrically controlled lens³¹. Combined with improved data acquisition technique, this method reduces the time necessary for a single scan from a few minutes to limiting several seconds, thus the possibility of photodegradation. While the majority published vast of papers described only the singlet oxygen photogeneration process, or two-photon absorption properties of the studied photosensitizers, in this work both are treated with equal attention and detail, allowing for more reliable conclusions.

The spectra of two-photon absorption cross-sections of investigated porphyrins obtained by f-scan technique are shown in Figure 3. They are overlaid with linear absorption spectra, shown as red curves, plotted *vs.* twice the wavelength. For all

porphyrins, both the monomer (1) and dimers (2-4) the maximum of two-photon absorption cross-sections is within first biological window which exists at wavelengths between 650 nm and 950 nm¹⁴.

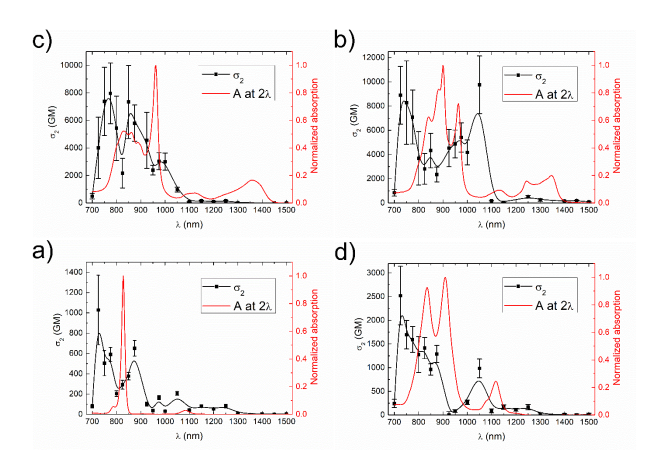


Figure 3. Spectra of two-photon absorption cross-sections of the investigated porphyrins (1-4). The spectra were overlaid with linear absorption (red curves) plotted vs. twice the wavelength. Figures a) to d) correspond to compounds 1 to 4.

In case of the monomer 1, the maximum means that it is favorable for the value of σ_2 is around 1000 GM at 725 nm, molecules to be excited to higher vibronic as shown in Figure 3. a). One can see that state upon 2PA excitation, compared to the band of 2PA is split into two, even 1PA process. Despite the fact that linear though the linear absorption shows only absorption of 4 is split into two equally one peak at 414 nm. It is caused by intense bands, the 2PA spectrum is different selection rules for one- and different. One can see that the band at two-photon absorption, as described in the 725 nm is much more intensive that that at literature²². It is worth noticing that the high 875 nm. The maximum value of σ_2 , energy 2PA band at 725 nm is observed measured at 725 nm is approximately for all the compounds and has the highest doubled compared to those of compound value of σ_2 among other 2PA bands. It 1 and is equal to 2500 GM. In the studies

of two-photon absorption cross-sections but its position remains the same, the various normalization factors are used, wavelength of maximum σ_2 is equal to 725 nm. However, the low energy band, among which the normalization to molar mass is the most popular. If the maximum which is located at 875 nm for other σ_2 is divided by molar mass, a similar investigated compounds, is value is obtained, which means that such bathochromically shifted to 1050 nm. structural change does not impose neither Similarly to compound 1, there is a dip in cooperation nor deleterious effects on the the two-photon absorption cross-section multiphoton absorption properties of the spectra at twice the wavelength of investigated compounds 1 and 4. In case maximum 1PA band. The 2PA spectrum of compounds 2 and 3, the spectra of of compound 3 is presented in Figure 3. c). two-photon absorption cross-section show The higher energy band is more complex structures, similarly to their bathochromically shifted by 50 nm to 1PA spectra. compound 2 775 nm and the value of σ_2 is around 8000 For (Figure 3. b), the higher energy 2PA band GM slightly being lower than for compound 2. For both compounds 2 and is broader than for compounds 1 and 4,

3, the maximum value of two-photon The values of maximum two-photon absorption cross-section is enhanced few absorption cross-section and the figure of times, while keeping the molar mass merit are listed in Table 2. The dimers of similar to compound 4. The results prove zinc porphyrins, especially connected by that a simple modification of the linker, ethyne and di-ethyne link, are much more from single bond (4) to ethyne (2) and dieffective two-photon photosensitizers than tetraphenyl porphyrin, or other porphyrin ethyne (3)links between the zinc porphyrins cooperation derivatives that are commercially used for cores cause effect, enhancing the σ_{2} * figure of merit. PDT.

Table 2. Two-photon absorption cross-section, merit factors and two-photon excited oxygen sensitization properties of the investigated porphyrins (1-4).

						Two-photon excited
	λ_{2PA}^{max}	σ_2^{max}	$\sigma_2^{max} *$	$\sigma_{2}^{max}\cdot\phi_{\Delta}$	$\sigma_{2}^{max}.\phi_{\Delta}*$	oxygen
	(nm)	(GM)	(GM·mol/g)	(GM)	(GM·mol/g)	sensitization
						enhancement factor ^[c]
1	725	1030	1.4	155	0.21	21
2	725	8900	6.1	3115	2.15	433
3	775	7950	5.4	3737	2.53	519
4	725	2500	1.7	675	0.45	94
H ₂ TPP	790	12 ^[a]	0.02	7.2 ^[b]	0.01	1

[[]a] taken from reference 42

[b] taken from reference 27

[c] Enhancement factor: $\sigma_2^{max} \cdot \phi_\Delta$ of the compound normalized to that of H₂TPP.

It was previously shown e.g. by TPEF conditions in the three-level system and dramatic enhancement of the excitedmeasurements¹⁷, that dimerization of porphyrin units leads to enhanced 2PA state singlet-singlet transition due to linear cross-sections and enhanced two-photon elongation of the π -conjugated system. production of singlet oxygen. The σ_2 Insertion of side groups, being either determined for the porphyrin dimer linked electron acceptors (A) or donors (D) by ethyne and di-ethyne bonds were on influence the σ_2 in a way that two-photon the order of 8200 and 5500 GM, which is absorption cross-section is the lowest for similar to the values found in the current higher neutral porphyrins, for the work for similar porphyrin derivatives. The porphyrins extended with A and the highest for compounds with D groups⁴³. cooperative effect was explained by a combination of several factors, including Even though the vast majority of molecules targeted to be used in 2PA PDT strong enhancement of the lowest onephoton Q-transition, better resonance are based on the porphyrin structure,

there are some alternative approaches with reasonable figures of merit. One example is macromolecular а photosensitizer, which was obtained via the covalent anchoring of several molecular PS based on а dibromobenzene (DBB) core onto a watersoluble polymer chain based on poly(N-acryloylmorpholine). It resulted in large 2PA cross-section up to 2600 GM at 740 nm and was proved to be an efficient candidate to cause melanoma cells death upon photo irradiation⁴⁴.

4. CONCLUSIONS

In this work, we investigated two-photon absorption and singlet oxygen

photogeneration properties in a series of zinc porphyrins, either in a monomer or dimer form. The dimers were connected by single bond, or by ethyne, or di-ethyne link. We observed strong enhancement of the obtained vields for quantum two-photon excited singlet oxygen generation (expressed by the product σ_{γ}^{max} the porphyrin dimer in connected by di-ethyne link, around 3700 GM, which is around 500 times more efficient than tetraphenyl porphyrin. It can be seen that in case of zinc porphyrin derivatives, structure optimization towards high two-photon absorption cross-section simultaneously enhances the properties of singlet oxygen photogeneration. Thus, in order to obtain a material capable of efficient production of singlet oxygen upon

increase the conjugation length, so that

excitation,

 π -delocalization is maximized, while

maintaining planar structure, as shown for

compounds 2 and 3. The results are very

promising for applications involving

treatment of deep tumors by

photodynamic therapy.

two-photon

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Conflict of interest statement

There are no conflicts to declare.

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