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# **Highly efficient and reproducible evanescent Raman converters based on a silica nanofber immersed in a liquid**

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### **Abstract**

We present highly efficient and reproducible Raman converters built with a silica nanofiber immersed in ethanol. The converters are pumped at 532 nm in the sub-nanosecond regime and the frst Stokes order photons are generated in the evanescent feld probing the liquid at 630 nm. Two standard fbers (SMF28 and 460HP) are tested and compared. The Raman conversion operating range limited by the damage threshold is optimized, leading to an external Raman conversion efficiency up to 60% with a nanofber radius of 300 nm and a length of 8 cm. The extracted Stokes energy is 0.29 μJ, which is three times higher than the previous result. We give guidelines for the design of other efficient evanescent Raman converters, opening the way for a new family of all-fbered compact Raman sources.

# **1 Introduction**

Optical tapered nanofbers are fabricated by pulling fbers until reaching diameters comparable or smaller than the light wavelength. Silica nanofbers are signifcantly exploited for a wide range of potential applications. Indeed, at such diameters, nanofbers exhibit a strong confnement of light which enables the generation of nonlinear efects [[1](#page-6-0)] such as, for example, the generation of supercontinuum [\[2\]](#page-6-1). Nanofbers can also exhibit an intense evanescent feld which has been exploited for optical sensing [\[3\]](#page-6-2), optical traps [[4](#page-6-3)] or

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spectroscopy [\[5\]](#page-6-4). However, the experimental investigation of the optical nonlinearities in the evanescent feld of the nanofber remains limited. Raman scattering is a third-order nonlinear efect that has been extremely widely studied in the literature in diferent media (solid, gas, and liquid) and different configurations (laser, fibers, free space,...) for example to generate new wavelengths [[6,](#page-6-5) [7](#page-6-6)]. In this study, we focus on the Raman scattering in the evanescent feld of a nanofber immersed in a liquid as the "evanescent nonlinearity". The Stokes photons are generated in the immersing liquid by the Raman scattering of the pump photons present in the evanescent feld and then couple to a guided mode of the nanofiber  $[8, 9]$  $[8, 9]$  $[8, 9]$  $[8, 9]$ . The observation of the evanescent Raman efect was previously investigated with a few-cm length nanofber immersed in ethanol and in toluene using sub-nanosecond pump pulses at 532 nm. This frst demonstration was performed with nanofbers having the maximal modal Raman gain [[9\]](#page-6-8). The output energy of the frst Stokes order of the Raman liquid was limited to 0.11 μJ by breakdown of the nanofber and never exceeded the transmitted pump energy.

To increase the Raman conversion operating range and to extract more Stokes energy, it is necessary to decrease the Raman threshold and to increase the optical nanofber breakdown threshold. For this latest condition, one way is to decrease the pump pulse laser duration possibly until a few tens of ps. Another way, keeping the sub-nanosecond regime, is to increase the diameter of the nanofber. However, higher diameters would increase the Raman threshold because the intensity of the evanescent field would be decreased. A compromise has to be found to determine the nanofber diameter enabling the extraction of the maximal output energy at the Stokes wavelength. In this paper, we study the design of the whole component (the nanofber waist, the tapers and, respectively, the input and the output untapered parts) not to only observe the evanescent Raman scattering routinely but also to extract the maximum of the Stokes energy. Two diferent fbers are compared. We defne the guidelines that lead to create an efficient  $\mu$ J Raman source in the sub-nanosecond regime.

The paper is structured as follows: we dedicate the frst section to detail the design of the component. The experimental setup is described in the second section. In the last section, the Raman conversion operating range versus the nanofber radius is depicted and the experimental performances of the optimized component are presented, showing an output Stokes energy of 0.29 μJ at 630 nm, i.e., a value that is almost three times higher than the previous results [[9\]](#page-6-8). We obtain external Raman conversion efficiencies around 60% with a high reproducibility. This opens the way to a new family of low-cost compact and efficient allfbered Raman sources that can be directly inserted in optical fbered networks.

### **2 Design of the component**

The component that we aim to design is presented in Fig. [1.](#page-1-0) The nanofiber is attached to two untapered fibers through two tapered sections called the tapers. The length of the nanofiber is  $L<sub>w</sub>$ . The two tapers are symmetrical, and their length is  $L_t$ . The length of the input (respectively, output) fiber is  $L_1$  (respectively,  $L_2$ ). The nanofiber and the tapers are totally immersed in the Raman liquid.  $L_1$  and  $L_2$  are required to make the manipulation easier and to facilitate the light injection and collection. The nanofber pulling rig is described in  $[9]$  $[9]$  $[9]$ . A butane flame softens the fiber central part while two computer-controlled translation stages

# $E_{p,out}$  $E_{i}$  $E_{s,out}$  $\boldsymbol{0}$  $\frac{1}{z}$

<span id="page-1-0"></span>**Fig. 1** The component to be designed

Y

elongate it following the "pull and brush" technique to create the nanofber and the tapers [\[10\]](#page-6-9). The pulling process is divided into cycles. Each cycle adds a small section to the taper while the radius of the fber cladding is reduced by a constant ratio to reach the nanofber waist radius at the end of the last cycle. At cycle iteration number *n*, the heat zone length is chosen as a power law of *n*, enabling to create fewcm length tapers.

This fabrication process is performed in a class-5 clean room. As soon as the nanofber is immersed in the Raman liquid, the experiment can be carried on in normal conditions. To size our component, we are based on the critical parameter  $\gamma$  expressed as  $\gamma = g_{\text{RM}} L_w P_{\text{in}}$  where  $g_{\text{RM}}$  is the modal Raman gain (in m<sup>-1</sup> W<sup>-1</sup>) and  $P_{in}$  is the peak pump power [\[11](#page-6-10)]. We consider that the Raman threshold is achieved when the transmitted pump power equals the output Stokes power, which corresponds to *γ*≈23 for sub-nanosecond pulses at  $\lambda_p$ =532 nm [[9\]](#page-6-8). The modal Raman gain  $g_{RM}$ is expressed following the analysis conducted in [[12\]](#page-6-11) by:

$$
g_{\rm RM} = \frac{\varepsilon_0 c^2 \iint_{\text{active area}} g_L n_L^2 \left| e_p \cdot e_s \right|^2 dA}{\iint_{\text{total area}} \left( e_p \times h_p \right) \cdot \hat{z} dA \iint_{\text{total area}} \left( e_s \times h_s \right) \cdot \hat{z} dA}.
$$
\n(1)

 $g_{\text{RM}}$  is depending on the liquid parameters which are the Raman gain coefficient of the liquid  $g_L$  (in m W<sup>-1</sup>) and its refractive index  $n_l \cdot \epsilon_0$  and *c* are, respectively, the dielectric permittivity and the light celerity in vacuum; *e* and *h* represent, respectively, the electric and the magnetic felds; p and s stand for, respectively, pump and Stokes. Due to the strong confnement of light and the high index contrast between silica ( $n = 1.45$ ) and the liquid ( $n<sub>L</sub> = 1.36$  for ethanol), the weak guidance approximation is no longer valid. We use a vectorial analysis to describe the modes propagation in the nanofiber waist. In the expression of  $g<sub>RM</sub>$ , the integral at the numerator is taken over the active area which is the liquid surrounding the nanofber and the denominator is a normalization factor. We consider that both the pump and the Stokes beams are propagating on the fundamental mode  $HE_{11}$ . Indeed, the modal Raman gain is very weak for crossed interactions between mode  $HE_{11}$  and higher order modes because of the small spatial overlap between the optical felds. As an example, the maximal modal Raman gain for the crossed interaction between mode  $HE_{11}$  and mode TM<sub>01</sub> is only 0.137 m<sup>-1</sup> W<sup>-1</sup>. We choose to use pure ethanol as the Raman liquid since it is nontoxic and easily manipulated. Its Raman gain coefficient is moderate (*g*<sup>L</sup> = 2.92 × 10<sup>−</sup><sup>12</sup> m W<sup>−</sup><sup>1</sup> at *λ*p=532 nm [\[13\]](#page-6-12)). The ethanol Raman shift is equal to 2928 cm<sup>-1</sup> at  $\lambda_p$ =532 nm and the first-order Stokes photons are generated at  $\lambda_s = 630$  nm. Figure [2](#page-2-0) shows the variation of the modal Raman gain  $g_{\rm RM}$ and the incident energy at the Raman threshold  $E_{\rm R, th}$  versus the nanofber radius R when the nanofber is immersed in



<span id="page-2-0"></span>**Fig. 2** Modal Raman gain (left axis) and Raman threshold energy (right axis) versus the radius of a nanofber immersed in ethanol. Empty square is for NF1, grey square is for NF2 and full black square is for NF3

<span id="page-2-1"></span>Table 1 Geometrical parameters of the nanofibers under test

Nanofiber under test	$L_{\omega}$ (cm)	$L_{\rm t}$ (cm)	$R$ (nm)	$g_{RM}$ (m <sup>-1</sup> W <sup>-1</sup> )
NF1	8	5	220	0.98
NF <sub>2</sub>	8	4.2	300	0.61
NF3		4	350	0.37

ethanol. The target waist length  $L_w$  is 8 cm, which is the maximal length allowed by our pulling system.  $E_{\text{R},th}$  is computed using the thumb of rule  $E_{\text{R},\text{th}} \approx P_{\text{in}}\Delta t$  ( $\Delta t = 900 \text{ ps is}$ the FWHM of our laser pulse duration and  $F=4.7$  kHz is its frequency repetition rate) and using the expression of *γ* to compute  $P_{\text{in}}$  at Raman threshold. As we can see on Fig. [2,](#page-2-0)  $E_{\text{R},\text{th}}$  is increasing with *R*. For a nanofiber radius  $R = 220$  nm,  $g_{RM}$  is maximal and  $E_{R,th}$  is minimal which would intuitively lead to the highest extracted Stokes energy. When *R* is smaller than this radius, the evanescent feld is spread along important distances and its amplitude decreases reducing the modal Raman gain. Larger radii make the mode more confned in the silica core and the Raman gain decreases.

The nanofbers under test are denoted by NF1, NF2 and NF3 and their geometrical parameters are summarized in Table [1](#page-2-1).

We compare two standard fibers from which the nanofibers are pulled: a single-mode fber at 1550 nm (ref. SMF28 from Corning) and a single-mode fiber at  $\lambda_p = 532$  nm and  $\lambda$ <sub>S</sub> = 630 nm (ref. 460HP from Thorlabs). The fibers parameters (core radius  $r_{\rm co}$ , cladding radius  $r_{\rm clad}$ , core index  $n_{\rm co}$ , and cladding index  $n_{\text{clad}}$  are summarized in Table [2.](#page-2-2)

For the SMF28 fber, as the fber is highly multimode at  $\lambda_p$ =532 nm, high-order modes are filtered by pulling  $L_1$ over 1 cm until reaching a core radius of 1.3 μm and a cladding radius of 20 μm allowing only the propagation of the fundamental mode. We checked that the light beam remains Gaussian over 20 cm after the single-mode filter.  $L_1$  and  $L_2$ are, respectively, equal to 30 cm and 40 cm. Such lengths are long enough for practical use but short enough to avoid parasitic nonlinear efects as we will see later.

To model the propagation of the modes in the tapers, we assume that the modes are scalar in this region. As the taper is also immersed in the liquid, we use a three-layer model with two interfaces, core/cladding and cladding/liquid [\[16](#page-6-13)]. The adiabaticity of the fabricated tapers is examined. An adiabatic taper requires that the tapering angle should be small enough to prevent the propagating mode coupling to other undesired order modes, leading to high light transmission of the considered mode. In the following, we will study tapers adiabatic for the fundamental mode  $LP_{01}$ . The condition for adiabatic propagation usually adopted is that everywhere [[17\]](#page-6-14):

<span id="page-2-3"></span>
$$
\frac{1}{Z_{12}} \ge \frac{1}{r} \frac{dr}{dz}.
$$
 (2)

 $r(z)$  is the local radius of the taper transitions (see Fig. [1](#page-1-0)), *r* d*r*  $\frac{dr}{dz}$  is the normalized taper slope,  $Z_{12} = \frac{2\pi}{\beta_1 - \beta_2}$  is the beating length.

 $\beta_1$  is the local propagation constant of the pump mode  $LP_{01}$  and  $\beta_2$  is the local propagation constant of the mode having the highest chance to couple with it because of the azimuthal dependence, i.e.,  $LP_{02}$ .

The condition  $\frac{1}{Z_{12}} = \frac{1}{r}$ d*r*  $\frac{d}{dz}$  provides an approximate boundary between adiabatic and non-adiabatic (or lossy) behavior.

In Fig. [3](#page-3-0)a–d, we have represented the adiabatic boundary curves for diferent situations. We have also plotted the normalized taper slope of the experimental taper we have used for NF2 (in dotted line). This taper profle was short and adiabatic enough for our experiments, but other profles could be used.

<span id="page-2-2"></span>**Table 2** Fibers parameters:  $n_{\text{clad}}$  is calculated using the Sellmeier equation of pure silica, and  $n_{\text{co}}$  is deduced from the numerical aperture values given by the constructer [[14](#page-6-15), [15\]](#page-6-16)

	$r_{\rm co}$ (µm)	$r_{\text{clad}}$ (µm)	$n_{\rm co}$ at 532 nm	$n_{\text{clad}}$ at 532 nm	$n_{\rm co}$ at 630 nm	$n_{\text{clad}}$ at 630 nm
SMF <sub>28</sub>	4.1	62.5	.4628	. 4569	.4629	.4571
460HP	l.25	62.5	.4637	. 4569	. 4638	.4571



<span id="page-3-0"></span>**Fig. 3** Adiabatic boundary curves calculated from Eq. [\(2\)](#page-2-3). Experimental taper slope in dotted line. **a** SMF28 taper immersed in air  $(n<sub>air</sub>=1, full round)$ , in ethanol  $(n<sub>ethanol</sub>=1.36, empty round)$ , in isopropanol  $(n_{\text{isopropanol}}=1.38$ , crossed round). **b** 460HP taper immersed in air (full round), in ethanol (empty round), in isopropanol (crossed



round). **c** SMF28 taper immersed in air (full round), in ethanol (empty round), 460HP taper immersed in air (full square), in ethanol (empty square). **d** Tapers immersed in ethanol at 532 nm for the SMF28 (empty round) and 460HP (empty square). Same at 630 nm for the SMF28 (cross) and 460HP (star)

The tapers should be adiabatic in the immersing liquid even if pulled in the air. This condition helps to relax the adiabaticity constraints and to fabricate shorter tapers of a few-cm length.

Indeed, for both fbers, increasing the external medium refractive index shifts up the adiabatic boundary curves as shown in Fig. [3a](#page-3-0), b showing that it is less constraining to be adiabatic in ethanol than in the air and less constraining to be adiabatic in isopropanol than in ethanol. This observation is confrmed experimentally. When the nanofber is immersed in ethanol, the light transmission of the whole component at 532 nm defned as the ratio between the output energy  $E_{p,out}$  and the input energy  $E_{in}$  systematically increases: it typically goes from 83% for the SMF28 to 92% and from 80 to 87% for the 460HP. Immersing the nanofber in isopropanol makes the light transmission of the whole component increase from 83 to 95% for the SMF28 and from 80 to 90% for the 460HP.

Making the taper adiabatic with SMF28 is easier than with 460HP. Figure [3c](#page-3-0) shows that with ethanol and for the SMF28 taper the adiabatic boundary curve has a punctual minimum of 0.0035  $\mu$ m<sup>-1</sup> for a relative radius of 0.15. For the 460HP taper, this minimum is not as punctual. Its value is 0.0019  $\mu$ m<sup>-1</sup> just slightly above the maximal experimental slope value and has to be maintained over a relative radius range extending from 0.3 to 0.55 making the taper very sensitive to small deviations.

We checked that the light transmission remains routinely high: around 90% for both SMF28 and 460HP nanofibers when immersed in ethanol.

We also pay attention to the tapers adiabaticity at  $\lambda_{\rm S}$ . As it has already been observed in [[18](#page-6-17)] for another fber (SM980 from Newport), the plots in Fig. [3d](#page-3-0) show that the adiabatic boundary curve is right-shifted when the wavelength increases so that the tapers will also be adiabatic at the Stokes wavelength of ethanol  $\lambda_s = 630$  nm.

### **3 Experimental setup**

Our experimental setup is shown in Fig. [4](#page-4-0). A frequencydoubled pulsed pump laser (from HORUS) emits a beam at  $\lambda_p$ =532 nm with a pulse duration of 900 ps (FWHM), a frequency repetition rate of 4.7 kHz, a maximum available peak pump power of 7 kW and a maximum pump energy of  $6.3 \mu$ J. A lens L with a focal length of 10 cm collimates the pump light and the association of a half plate and a polarizing beam splitter enables to control the incident pump energy since the laser is linearly polarized. Part of the pump beam is then refected by the two



<span id="page-4-0"></span>**Fig. 4** Experimental setup scheme for the optical characterization of evanescent Raman converters. *PM* powermeter. PM1 is used for the reference measurement of the incident pump power. PM2 (resp. PM3) is used for the measurement of the backward (resp. forward) powers. *OSA* optical spectrum analyzer, *IR flter* infrared flter, *PBS* polarizing beam splitter. M1 and M2 are alignment mirrors

mirrors M1 and M2 to pass through a glass plate. The glass plate enables to analyze the backward beams. The pump beam is then injected by a microscope objective (x20, NA 0.35) in the untapered input end of the fber, propagates in the nanofber immersed in the liquid and is collected at the output end of the fber. The forward beams are analyzed thanks to the optical spectrum analyzer OSA1. The immersing tank length is about 20 cm remaining higher that the whole nanofiber length  $(L_w + 2L_t)$  for all treated cases, which guarantees the total immersion of the waist and the tapers in the Raman liquid.

We are first interested in the three nanofibers SMF28-NF1, SMF28-NF2, and SMF28-NF3. To optimize the evanescent Raman conversion, avoiding the undesired backward and forward nonlinear efects is mandatory. We checked that  $L_1$  and  $L_2$  are short enough to avoid nonlinear efects in the fber and we investigate nonlinear efects in the nanofber immersed in ethanol. We present in Fig. [5](#page-4-1) the forward output spectrums for the three nanofbers immersed in ethanol just before they break. We checked that there is no signal at 545 nm which is the Stokes silica frst-order wavelength. We also checked that there is no backward silica Raman scattering by visualizing the backward energy refected by the glass plate with OSA2. We fnally measure the backward energy at 532 nm as a function of the laser incident energy for each nanofber. As the backward energy increased linearly with the laser incident energy, we conclude that there is no nonlinear backward scattering. The backward energy was very weak and attributed to parasitic refections.



<span id="page-4-1"></span>**Fig. 5** Collected forward spectra from OSA1 for SMF28-NF1 (full line), SMF28-NF2 (dotted line) and SMF28-NF3 (dashed line) immersed in ethanol just before they break

#### **4 Results**

Since undesired forward and backward beams are avoided, we focus on the measurements of the maximum extracted Stokes energy with using ethanol as an immersing liquid. We test the reproducibility of the experimental performances by drawing and manipulating 20 samples of each nanofber (SMF28-NF1, SMF28-NF2, SMF28-NF3). As an example, we plot in Fig. [6](#page-4-2) the experimental Raman threshold energies, incident nanofber breakdown energies and maximum extracted Stokes energies for the 20 samples of SMF28- NF2 underlying the reproducibility of the performances. The same results in term of reproducibility (not shown here) are obtained for NF1 and NF[3](#page-5-0). In Table 3, we compute the average and the standard deviation of the maximal extracted Stokes energy values  $E_{\text{Smax}}$  for the 20 samples of each nanofber just before the breakdown.



<span id="page-4-2"></span>**Fig. 6** Experimental Raman threshold energies (triangles), incident nanofber breakdown energies (rounds) and maximal Stokes energies (squares) for the 20 samples of SMF28-NF2

<span id="page-5-0"></span>**Table 3** Average and standard deviation of the maximal extracted Stokes energy values  $E_{\text{Smax}}$  computed for the 20 samples of each nanofber just before breakdown

Nanofiber	Average of $E_{\text{Smax}}$ ( $\mu$ J)	Standard devia- tion of $E_{\text{Smax}}$ $(\mu J)$	
NF1	0.126	0.007	
N <sub>F2</sub>	0.283	0.007	
NF3	0.147	0.006	

Figure [7](#page-5-1) shows typical results. The transmitted pump energy  $E_{p, \text{out}}$  and the output Stokes energy  $E_{s, \text{out}}$  are plotted as a function of the incident pump energy  $E_{\text{in}}$  for the three nanofbers (SMF28-NF1, SMF28-NF2, SMF28-NF3). For low energies, the transmitted pump energy increases linearly and begins to be depleted when the Stokes photons are generated. We define the external efficiency of the Raman conversion  $\eta_{ex}$  as the fraction between the maximum output Stokes energy and the incident pump energy after reaching the Raman threshold. The highest Stokes energy was obtained with SMF28-NF2. We reach an energy of 0.29 μJ at 630 nm just before the SMF28-NF2 breaks for  $E_{\text{in}} = 0.48 \mu\text{J}$ , which provides  $\eta_{ex}=60\%$ . The Raman threshold is reached at  $E_{\text{in}}$  = 0.41 μJ. For SMF28-NF1 which is the nanofiber having the highest modal Raman gain, the maximum extracted Stokes energy is only 0.13  $\mu$ J and  $\eta_{ex}$  is 51%. The Raman threshold is directly followed by the breakdown of the nanofber. With SMF28-NF3, we are collecting a Stokes energy of only 0.15  $\mu$ J for an incident energy  $E_{in}=0.58 \mu$ J and the Raman threshold is not reached.

To figure out the reason behind not obtaining the highest extracted Stokes energy with SMF28-NF1 as it could be intuitively expected, we plot in Fig. [8](#page-5-2) the experimental Raman threshold energy and the experimental incident nanofber breakdown energy versus the radius of the nanofber when the nanofber is immersed in ethanol. These two plots defne the boundaries of the Raman



<span id="page-5-1"></span>**Fig. 7** Transmitted pump energy  $E_{p,out}$  at 532 nm and output Stokes energy  $E_{s, \text{out}}$  at 630 nm versus the laser incident energy  $E_{\text{in}}$  for SMF28-NF1, SMF28-NF2 and SMF28-NF3



<span id="page-5-2"></span>**Fig. 8** Experimental Raman conversion operating range and theoretical Raman threshold boundary versus *R*. The experimental Raman threshold energy at  $R = 350$  nm (NF3) is extrapolated from Fig. [7](#page-5-1)

conversion operating range  $\Delta E_{\rm R,\,break}^{\rm exp}$ . We also plot the theoretical Raman threshold energy deduced from the expression of *γ*. The experimental and the theoretical Raman thresholds for NF1 and NF2 are in a good agreement and confrm the value of the critical parameter *γ*≈23 obtained in [[8](#page-6-7)] which is higher than the value of 16 presented by [[11\]](#page-6-10). This discrepancy is explained by the higher gain and shorter length we used in nanofiber experiment, compared with the standard fibers studied in [[11\]](#page-6-10). Based on a linear extrapolation from Fig. [7,](#page-5-1) we can estimate the Raman threshold would be obtained for  $E_{\text{in}}=0.6 \mu \text{J}$  for NF3 if the breakdown was not obtained before. Figure [8](#page-5-2) shows that there is a maximal Raman conversion operating range for a radius of  $R = 300$  nm. At this radius,  $\Delta E_{\text{R, break}}^{\text{exp}} = 0.08 \,\mu\text{J}.$ When the radii are below 300 nm, the experimental Raman conversion operating range decreases. It is divided by 2 for NF1  $(R = 220 \text{ nm})$  which explains the inability of producing the maximum of Stokes energy even if the nanofiber has the highest modal Raman gain  $g_{RM}$ . For radii higher than 300 nm,  $\Delta E_{\text{R, break}}^{\text{exp}}$  also decreases and we are even no longer able to experimentally reach the Raman threshold starting from 340 nm, as confrmed by the "closure" of the Raman conversion operating curve.

Since the nanofber breakdown limits the performances of the component, we compare in Table [4](#page-6-18) the incident breakdown energies  $E_{\text{break}}$  of the SMF28 nanofibers surrounded, respectively, by air and ethanol. Almost the same values are found with 460HP nanofbers. This simple criterion gives already interesting trends. In air, the breakdown energy increases with the radius. When the nanofber is immersed in ethanol, the breakdown energy becomes higher for all the considered radii. The breakdown of a nanofber immersed in a liquid in the sub-nanosecond regime is a complex mechanism with a thermal origin and

<span id="page-6-18"></span>



depends on the fraction of the light in the silica and in the liquid. Depending on the radius of the nanofber, the surrounding medium refractive index and the thermal conductivities of the liquid and silica, the breakdown can be induced either in the silica or by cavitation bubbles in the liquid. A complete study of the breakdown of immersed and non-immersed nanofbers induced by light will be published elsewhere.

## **5 Discussion and conclusion**

We have demonstrated and optimized evanescent Raman converters in the sub-nanosecond regime based on a silica nanofber immersed in ethanol. The converters were pumped at 532 nm and delivered pulses at 630 nm, which is the wavelength of the frst Stokes order of ethanol. Two diferent standard silica fbers were tested and compared (SMF28 and 460HP). These evanescent Raman converters are lowcost alternatives to Raman converters based on hollow core photonic crystal fbers (HCPCF) flled with liquids [\[19](#page-6-19), [20](#page-6-20)]. The two technologies present Raman conversion efficiencies of the same order of magnitude. Some diferences can be underlined: with nanofbers, there is no possibility to control the eventual Raman cascade, which was the case using liquid-flled HCPCF. Another diference is that we observed no Brillouin backward scattering from the liquid. On the contrary, this had to be considered in the design of liquid-flled HCPCF since this efect was in competition with Raman scattering  $[21]$ . This absence is probably due to the fact that Brillouin is generated at the maximum of the pump feld, which is located inside the silica for nanofbers and not in the immersing liquid [\[22\]](#page-6-22).

The performances of the evanescent Raman converters are highly reproducible. We obtained a maximum output Stokes energy of  $0.29 \mu J$  with an external conversion efficiency from the pump to the frst Stokes order of ethanol of  $60\%$ . This conversion efficiency is limited by the Gaussian temporal pump pulse, as it has been shown in [[19\]](#page-6-19). To optimize the extracted Stokes energy, we have defned for each nanofber radius an operating range, whose boundaries are on one side the Raman threshold energy and on the other side the breakdown energy. In a way that is counter intuitive, the optimized nanofber had a radius which does not correspond to the maximum of the Raman gain. This operating range is a guideline for the conception of other evanescent Raman converters based on other liquids and/or fbers. Other perspectives are foreseen such as the deposition of nonlinear polymers on the nanofber surface opening the way to a new family of robust all-fbered components that can be directly inserted in optical networks without additional losses.

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