Fiber-Optic Temperature and Pressure Sensors Applied to Radiofrequency Thermal Ablation in Liver Phantom: Methodology and Experimental Measurements

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Received 14 October 2014; Accepted 15 December 2014

Academic Editor: Geoffrey A. Cranch

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Radiofrequency thermal ablation (RFA) is a procedure aimed at interventional cancer care and is applied to the treatment of small- and midsize tumors in lung, kidney, liver, and other tissues. RFA generates a selective high-temperature field in the tissue; temperature values and their persistency are directly related to the mortality rate of tumor cells. Temperature measurement in up to 3–5 points, using electrical thermocouples, belongs to the present clinical practice of RFA and is the foundation of a physical model of the ablation process. Fiber-optic sensors allow extending the detection of biophysical parameters to a vast plurality of sensing points, using miniature and noninvasive technologies that do not alter the RFA pattern. This work addresses the methodology for optical measurement of temperature distribution and pressure using four different fiber-optic technologies: fiber Bragg gratings (FBGs), linearly chirped FBGs (LCFBGs), Rayleigh scattering-based distributed temperature system (DTS), and extrinsic Fabry-Perot interferometry (EFPI). For each instrument, methodology for ex vivo sensing, as well as experimental results, is reported, leading to the application of fiber-optic technologies in vivo. The possibility of using a fiber-optic sensor network, in conjunction with a suitable ablation device, can enable smart ablation procedure whereas ablation parameters are dynamically changed.

1. Introduction

Radiofrequency thermal ablation (RFA) is an emerging medical procedure for targeted treatment of tumors [1–6]. Pioneered through the ’90s by Rossi et al. [7, 8], RFA makes use of electrical irradiance through a miniature applicator percutaneously inserted at the point of treatment. Tumor cells are damaged by high-temperature conditions, and cytotoxicity is function of the local temperature value and its persistency [9]; 52°C for 60 s exposure is considered the clinical reference condition for cancer cells mortality, while over 60°C cells death is nearly instantaneous. Thermal dosimetry [8, 9] defines the temperature dose that is related to each mortality rate of tumor cells.

RFA makes use of RF irradiance to provide a selective heating, confined within the tumor [1, 8, 10]. A midpower (5–50 W) RF source operating at 350–500 kHz is used to power a miniature ablation device (1–4 mm diameter), percutaneously inserted in the tumor and with an electrical active region of up to 1 cm length; ablation devices having stem, multitiled, or other shapes have been reported [11, 12]. By powering the ablator with RF, the electrical irradiance propagates in the biological tissue by conduction. RFA generates steep spatial (3–5°C/mm) and temporal (0.4°C/s) gradients that ensure that the treatment is confined to the tumor.

The outcome of RFA depends on the evolution of the electrical impedance throughout the ablation. In most tissues, the impedance at ambient temperature is 20–200 Ω and slightly reduces during the first part of ablation. Whereas temperature at the ablation point reaches 95–100°C, the water constituents of the tissues reach the ebullience point; the impedance of vapor, at RF, is orders of magnitude higher than that in liquid water; hence, after reaching the ablation peak, the target tissue is progressively insulated from the electrically active area,
substantially neutralizing the irradiance. This phenomenon is extremely significant in hepatic tumors \cite{8, 9, 13–16}, as liver is a soft tissue with high water concentration. When RFA is applied to liver tumors, the vapor-to-liquid transition causes an instantaneous rise of the tissue impedance that abruptly interrupts the ablation procedure: this effect limits RFA to small hepatic tumors (up to 3 cm in size), while the treatment of mid-to-large tumors via percutaneous RFA is subject to disruptive improvements of the whole ablation setup.

The RFA procedure can be substantially improved by using sensors operating at the point of treatment. Currently, temperature measurement in RFA is performed with miniature thermocouples (TCs) \cite{15, 21, 22}. TCs allow measurement in up to 3–5 points within the tumor volume and can be either installed on the ablator or externally inserted. However, TCs are invasive and can perceivably alter the ablation pattern, suffer from hysteresis, and are limited to few sensing points. Alternatively to TCs, noncontact techniques such as infrared thermal imaging have been reported \cite{23, 24}, but they are limited to surface imaging with a direct line of sight.

Optical fiber sensors (OFS) \cite{25} can significantly improve thermal measurement in RFA. OFS are smaller than their electrical counterparts, so as not to alter the ablation pattern and exhibit hysteresis. The use of distributed or quasidistributed sensing techniques allows unmatched density of sensing points per units of length, providing sensing below 0.1 mm spatial resolution. The application of fiber-optic thermal sensors has been limited to few preliminary studies in RF \cite{17, 18, 26} and laser \cite{27, 28} ablation. Using OFS, it is also possible to increase the number of sensing points per unit of volume, by using inline wavelength division multiplexing (WDM) approaches and progressing towards quasidistributed and distributed sensing.

In this paper, methodologies for temperature detection in RFA are detailed, progressing from a fiber Bragg grating (FBG) array (1 sensing point/cm) to a linearly chirped FBG (LCFBG) for quasidistributed measurement (130 sensing points/cm) to a distributed temperature sensing system based on Rayleigh backscattering (100–500 sensing points/cm). Experimental measurements have been carried out \textit{ex vivo} on liver phantom; several chambers have been developed in order to confine the tissue and position the fiber along different ablation axes. In addition to temperature measurement, the pressure readout is a key asset to support the advanced treatment of encapsulated tumors \cite{29}. This measurement is addressed with a Fabry-Perot fiber-optic sensor \cite{18, 30}, with a low thermal coefficient, and is combined with a fiber Bragg grating sensor.

Biophysical sensing applied to RFA allows monitoring this treatment with unprecedented precision in real time. When RFA is applied to small ablations, for example, for thermal treatment of small tumors in hard tissues \cite{1, 2}, in correction of cardiac arrhythmia \cite{31} or for interventional pain management, it is possible to adequately control the ablation procedure by just timing the RF irradiance. However, when RFA is applied to the treatment of midlarge tumors, particularly in soft tissues, the ablation pattern has a strong dependence on the structure of the biological tissue, which often incorporates blood vessels or fibrotic capsules and therefore results in asymmetrical ablations, which can potentially fail to reach the entire tumor. The deployment of a sensor network can potentially be used to trigger a close-loop feedback to the ablation, changing the parameters of the RF source and of the ablation device as a function of sensors readout. Such concept, namely, smart ablation, has been only partially initiated with TC sensors \cite{21} but can be disrupted with OFS technologies.

The paper is arranged as follows: Section 2 describes the RFA setup, procedure, and measurement methodology; Section 3 outlines the measurement setup and the experimental results of Bragg grating-based measurements (FBG and LCFBG); Section 4 outlines the DTS setup for distributed measurement along multiple axes; Section 5 describes the pressure measurement and its significance; Section 6 discusses the measurement setups and their potential use in a smart RFA concept; finally, Section 7 draws conclusions.

2. RFTA Setup

Figure 1 illustrates the setup used for RFTA. The RF source is a medical-grade generator (TAG 100 W, Invatec), with 480 kHz sinusoidal carrier. The emission power is kept constant throughout the experiments, within the 9–20 W range in order to adjust the duration of ablation and the power density per unit of volume at the ablation point; most experiments have been carried out using 20 W electrical power. The RF generator embeds an impedance meter, which records the electrical impedance of the tissue in real time and disconnects the RF power when it falls outside the range 20–300 Ω.

Both voltage and current provided by the RF generator are recorded with an oscilloscope (Agilent Technologies DSO3062A, 60 MHz), in order to monitor the RF power input to the ablator.

RFTA experiments have been performed on samples of porcine liver, the natural phantom of the human tissue. In agreement with EU protocols, all specimens originated from pigs slaughtered no more than 48 hours in advance of the experiment and are commercially available; samples have been refrigerated until the ablation. The impedance of the tissues chosen for experiments is 60–120 Ω at ambient temperature; during the heating process, impedance slowly decreases (about 15%) until temperature reaches a plateau around ~95°C. Then, the water constituents of the liver transition from liquid to vapor state, and, as a consequence, the tissue impedance abruptly rises to several kΩ, causing the disconnection of the RF power and the interruption of the RFTA. This phenomenon is particularly evident in hepatic ablations due to the high concentration of watery constituents in the tissue, which cause an almost instantaneous rise of the electrical impedance in excess of one order of magnitude.

In some experiment, the liver sample has been positioned in a chamber built with extruded polystyrene (XPS) material, in order to confine the tissue in a finite volume preventing its expansion, as in Figure 1; other experiments have been designed leaving the explanted tissue external to the chamber.

The ablation device used in most experiments is a hollow brass stem-shaped needle with 3 mm inner and 4 mm outer diameter. The needle sustains the penetration in the tissue
Figure 1: RFTA setup. A radiofrequency generator (RFG) is used to power an ablation device having 3/4 mm inner/outer diameter and with 1 cm active length on the tip; the active electrode (AE) is connected to the ablator, while the passive electrode (PE) is connected to a plate below the measurement chamber, in a neutral spot. RF power is monitored in real time through an oscilloscope. The measurement chamber, made of extruded polystyrene (XPS), is designed in accordance to each experiment and internally hosts the porcine liver tissue. Two thermocouples (TCs) are used as reference, one positioned internally to the ablator, one positioned externally in the tissue, and both controlled with a data acquisition (DAQ) system in LabVIEW.

and has an electrically active electrode with 10 mm length on its tip, providing the electrical connection with the tissue. The needle is connected to the RF generator, while a plate positioned below the chamber is the second electrode; the difference of potential is therefore applied between the needle tip and the reference plate. In order to provide a temperature reference, a TC (0.1°C accuracy) has been positioned within the hollow ablation device, in correspondence of the center of the electrode.

3. Fiber Bragg Grating-Based Sensors

3.1. Fiber Bragg Gratings. FBG arrays represent the most straightforward technique for inline temperature measurement with fiber-optic technology [32–35]. By using an FBG array fabricated on a drawing tower [36, 37], it is possible to reduce the distance between adjacent sensors and building a sensor array with constant spacing between each active region. This allows an unambiguous labeling of the sensors at the point of treatment.

The setup for FBG sensing, illustrated in Figure 2, makes use of an array of 5 draw-tower FBGs (FBGS DTG-1550) fabricated on ormoceramic bend-insensitive fiber. Each active length is 5 mm, interleaved by 5 mm void, obtaining a sensing structure with 1 sensor/cm spatial resolution. FBGs have Bragg wavelengths around 1550 nm, with 1.8 nm spacing between adjacent FBGs, and 11.66 pm/°C thermo-optic coefficient [26]. Sensors are interrogated with a white-light setup, based on a broadband source (Optolink ASE, 20 mW, C + L bandwidths) coupled to a spectrometer (Ibsen I-MON-USB) for detection. The use of a grating array fabricated on a drawing tower, whereas the gratings have both uniform spacing in the spectrum, with identical wavelength separation at constant temperature and along the longitudinal axis of the fiber, allows a straightforward mapping of each measurement point.

Accurate demodulation of the FBG array is performed in the spectrum domain. In first place, the pitch wavelength of each FBG is tracked by means of a spectral correlation technique as in [38], with 0.07°C typical accuracy on each of the 5 measurement points. Then, using a spline function, the spatial resolution is artificially expanded from 1 cm to 1 mm by interpolating the inner measurement points. Figure 2(b) shows the spectrum of the FBG array, as recorded by the spectrometer, through a sample RFA procedure.

Experiments with the FBG arrays have been performed with two setups, photographically shown in Figure 3. The aim of this approach is estimating the axial temperature distribution (i.e., along the direction of the ablation device), whereas the thermal gradient is lower [15, 26]. The first setup mimics the insertion of the FBG array on the ablation device, as in Figure 3(a). In this setup, the ablation peak is located in correspondence to FBG4 (as labeled in Figure 2(a)). The second setup is a better representation of the FBG array inserted externally through the tissue, and the ablation peak is located within FBG2 and FBG3. All experiments have been carried out using a 4 mm hollow ablation device and 20 W RF power.

Figure 4(a) shows the result of the ablation performed with the first setup. After the initialization, temperature...
Figure 3: Measurement chambers used for FBG sensing in RFA, mimicking the FBG array embodied in the ablation device ((a)-(b)) and an external insertion (c). In the first setup, the FBG array (a) is attached to a 4 mm ablation device, stem-shaped, patching the fiber outside of the active electrode; the ablator is then inserted (b) in porcine liver lying on a metal plate serving as passive electrode. In the second setup (c), an extruded polystyrene chamber (10 × 8 × 6 cm) is set up, confining the liver tissue. A plexiglass layer encloses the chamber from the top, having the FBG array attached on its bottom side. A red laser is used to illuminate the fiber tip.

Figure 4: Temperature recorded by the 5 FBG sensors throughout two RFA procedures. (a) RFA performed using the setup as in Figure 3(a). (b) RFA performed using the setup as in Figure 3(c).
increases more than linearly in the first part of the ablation, reaching the 60°C threshold [9] after 51 s at the ablation peak (FBG4) and 111 s for FBG3. In a second phase, temperature grows almost linearly, with an approximate gradient of 0.35°C/s for FBG4 and 0.26°C/s for FBG3 within the 80–240 s interval. A final temperature rise is recorded, and temperature exceeds 100°C; at this stage, the rapid impedance increase of the tissue causes the RF generator to discontinue the power supply, and temperature starts to decrease. The temperature recorded by FBG1 and FBG2 is substantially lower [14, 15], as a reflection of the thermal gradient. Both charts show a similar ascending trend, with an extended linear range; FBG2 records a maximum temperature of 80.2°C and intersects the 60°C level after 255 s, while FBG1 does not record a cytotoxic temperature.

Figure 4(b) shows the RFA experiment with the setup as in Figure 3(c). In this case, the ablation peak lies between FBG2 and FBG3. FBG3 records the highest temperature (103°C); unlike the previous experiment, it exhibits a nonlinear growth in proximity of the ablation peak, which typical with the variability of RFA procedures. The chart shows that the final rise of temperature is experienced by FBG3 earlier than FBG2, with a delay of about 35 s. FBG1 and FBG4, positioned almost symmetrically with respect to the ablation peak, record a very similar pattern, and the temperature recorded by these sensors just overcomes the 60°C threshold in correspondence to the ablation peaks.

Ablation data can be reported as a thermal map, after upsampling temperature on the longitudinal axis. Such format, as in Figure 5, allows also a visual comparison between different ablation experiments. Contours in particular show the temperature pattern, which is approximately quadratic in the first experiments, almost linear in the second experiment, while, in the third RFA, the linear trend appears to be altered in proximity of FBG4.

Time and temperature normalization allow a quantitative comparison between different RFA experiments. The normalization takes into account the start of the RFA procedure and the final instant of the heating phase, before the RF power is disconnected; these initial and final values of temperature are normalized between the [0–1] range. The same normalization will be used for all the experiments.

The use of FBG sensing can also enable RFA prediction features [22]. Figure 6 reports the results of the same ablation experiments as in Figure 5, and both time and temperature have been normalized. In the second ablation procedure, which substantially differs from the other experiments as it has a longer duration, it is possible to notice that the FBG positioned at the peak (FBG2) records a lower normalized temperature as compared with the other ablations. On the other side, for all the other FBGs, the normalized temperature recorded in the second experiment is higher, with a significant deviation recorded in FBG3. Data recorded in Experiments 1 and 3, which exhibit a similar behavior, are nearly overlapping, with the exception of a difference recorded in proximity of FBG4. The possibility to use the thermal data recorded by the FBG array as a metric for the effect of RF irradiance in the tissue can be exploited as a predictive asset: the thermal distribution discriminates between a heat distribution that has a deeper penetration in the tissue, as in the first and last experiment, and conditions in which the temperature field is more confined near the ablation focus, as in the second experiment, which results in a smaller ablation volume.

3.2. Linearly Chirped Fiber Bragg Gratings. While uniform FBGs are a discrete array of sensors, with a tight identification of active and passive regions, chirped FBGs [32, 34, 39–41] allow extending FBG sensing to quasidistributed detection. A chirped FBG can be modeled as a cascade of reflectors [34, 42], each having a different Bragg wavelength; when a temperature variation is experienced in each part of the grating, the corresponding local pitch wavelength shifts linearly as for uniform FBGs [17]. By using a linear–chirp pattern, thus fabricating a LCFBG [17, 39, 43], it is possible to have a linear relationship between each grating portion and the correspondent local temperature variation. This allows using the LCFBG as a distributed sensor having an active region corresponding to the grating length, and a spatial resolution proportional to the wavelength resolution of the spectrometer.

The setup for LCFBG interrogation is illustrated in Figure 7. The optical equipment is the same used for FBG interrogation (Section 3.1). The sensor is a LCFBG (Technica S.A.) fabricated on standard single-mode fiber and with a 250μm protective coating; Figure 8 shows the grating spectrum. The grating length is 1.5 cm, while the 3 dB bandwidth is 33.4 nm (1529.3–1562.7 nm), which corresponds to a chirp coefficient of 2.22 nm/mm. The LCFBG has a maximum reflectivity of 94.5% and 2.0 dB ripple. When uniform temperature is applied, the LCFBG spectrum exhibits a thermooptic coefficient of 10.2 pm/°C.

The decoding of the whole temperature distribution through the LCFBG involves signal processing. Skaar’s reverse layer peeling [42, 44] is a popular method for grating synthesis but involves overwhelming computational...
Figure 6: Thermal pattern of RFA, considering only temperature rise. Both time and temperature variation from the initial offset are normalized within the [0-1] range. Data are reported for each FBG sensor, for the three RFA experiments as in Figure 5.

Figure 7: Setup for interrogation and decoding of the LCFBG. Interrogation is based on a white-light scheme, with a 1550 nm spectrometer for detection. The simplified decoding algorithm, as in [17], is initialized by dividing the measured LCFBG spectrum by the reference spectrum, acquired in reference condition (uniform temperature). The obtained spectral function is an almost flat curve, which depends on the whole temperature pattern, while the fluctuations at the two sides (the uppermost red and blue parts of the LCFBG spectrum) identify the temperature values at the two sides of the grating, that is, the “left” and “right” temperatures. The inner part of the spectrum is integrated and normalized for a calibration function acquired in reference condition, to estimate the thermal slope. Finally, each contribution is combined to obtain the temperature distribution through the whole LCFBG active length.

Figure 8: LCFBG spectrum acquired at ambient temperature (blue) and after 76 s (black) and 128 s (red) elapsed time in RFA heating.

in real time, Tosi et al. recently proposed a method based on spectral analysis, which provides a simplified detection based on simple spectral operations; this approach is sketched in Figure 7 and reported in [17] and returns the temperature with nonuniform accuracy of $0.1-2.5^\circ C$ along the grating length (from the head to the tail of the sensor) and a spatial resolution of $\sim 75 \mu m$ correspondent to 166 pm resolution of the spectrometer divided by the chirp coefficient.

Since the LCFBG has a relatively short length and the capability for a narrow spatial resolving, it has been used to monitor the ablation in the direction perpendicular to the ablation axis (hence, perpendicular to the FBG array sensor as in Section 3.1), whereas the depth of penetration of heat into the tissue is lower. As for the previous characterization, two measurement setups have been used, as in Figure 9. The first setup, similar to Figures 3(a)-3(b), is based on direct insertion of the ablation device into the liver phantom; the LCFBG is
Figure 9: Measurement setups used for RFA with LCFBG monitoring of the temperature along the perpendicular axis to the ablation device. The LCFBG is illuminated with a red laser, highlighting the whole active part where light scatters in the outer buffer. (a) RFA has been performed on plain liver, with a 4 mm ablation device (1 cm active electrode). The LCFBG is positioned perpendicularly to the ablation device at the center of the electrode. The ablation device is inserted through the tissue; the picture shows the liver phantom after the RFA treatment, cut in half to show the ablation pattern. (b) Porcine liver is confined in a cylindrical XPS chamber, with diameter 6 cm. The ablation device, same as (a), is inserted perpendicularly to the chamber plane from the bottom side. The LCFBG lies on the bottom plane, in contact with the tip of the electrode. The picture shows the chamber setup prior to the insertion of the liver phantom. RFA experiments have been performed using RF power of 20 W.

Figure 10: Thermal maps of RFA experiments, reporting the instantaneous temperature (color bar, in °C units) as a function of distance from the sensor tip and elapsed ablation time. (a) Setup as in Figure 9(a); (b) setup as in Figure 9(b).

positioned perpendicularly to the direction of ablation, in contact with the active electrode. The second setup, closer to Figure 3(c), encloses the porcine liver in a cylindrical chamber; ablation is performed on the bottom plane of the chamber, and the LCFBG sensor is fixed perpendicularly to the ablation device on the bottom plane, in contact with the electrode.

Figure 10 shows the results of two RFA experiments, carried out with the setups as in Figure 9; temperature is reported as a function of distance (measured from the LCFBG tip, positioned in contact with the electrode) and time. The qualitative analysis of the thermal maps shows similarities with the measurements recorded in Figure 4. The first chart shows an almost linear temperature trend, similar to Figure 4(a) and Experiments 1 and 3 in Figure 5. The narrow spatial resolution of the LCFBG highlights that the approximate linear trend occurs both in space and time, with approximate gradient 2.8°C/mm. This linear trend, as confirmed by the previous measurement, confirms that the depth of penetration of heat through the tissue is higher:
a temperature higher than 55°C is obtained through all the distances up to 12 mm before the RFA reaches the final abrupt temperature rise. On the other hand, the second chart shows a trend more similar to Figure 4(b): temperature has an almost quadratic rise over time, particularly close to the ablation peak, and the spatial distribution is nonlinear, with a perceivable distortion with respect to Figure 4(a). Visually, the thermal map appears almost flat along the distance axis, particularly within the 0.2–2.6 mm region. The 55°C threshold is achieved only for distances lower than 11.2 mm, which results in a lower effectiveness for tumor treatments. Figure 10, in line with [15, 22, 45], highlights that the ablation pattern has a strong variability, and it depends on the measurement chamber setup.

In order to emphasize quantitative results, Figure II reports the contour plot as in Figure 10(a). For short distances (0–0.6 mm), temperature is rapidly growing reaching a maximum of 104°C within 0.08 mm from the peak point. The middistance (2–5 mm) range entirely overcomes the 60°C threshold after 88 s well before the end of the RFA treatment. The thermal gradients in this range are nearly linear and almost constant. The long-range (10–14 mm) plot shows that the 60°C value is reached only for distance lower than 10.4 mm. This region defines the margin of ablation, and the accurate measurement of temperature can define the mortality rate of cancer cells [9].

4. Distributed Temperature Sensing

Distributed sensing provides an alternative approach over grating-based sensors [46–50]. DTS systems turn an entire fiber cable into a fiber sensor, without discrimination between active and passive regions. Optical time-domain reflectometry is a typical approach for achieving long-distance sensing [46–48], over multi-km range; on the other side, optical frequency-domain reflectometry (OFDR) [51–54] provides a technique for distributed sensing with a narrow spatial resolution.

The OFDR method proposed by Froggatt et al. [51–53], based on coherent demodulation of Rayleigh backscattering, allows achieving narrow spatial resolution, as low as 20 µm, over fiber length higher than 20 m [52]. The principle of operation is based on swept-wavelength interferometry (SWI) [53]. A tunable laser is used as light source, splitting the laser beam through a coupler between a measurement arm (connected to the fiber output to the DTS) and a reference arm (internal to the DTS unit); a second coupler recombines the two arms. Light is split into two orthogonal polarization states by means of a polarization controller between the two arms [50, 53] and a polarization beam splitter on the reference end. The SWI setup records the chaotic Rayleigh backscattering pattern, and, by means of the polarization-based setup, it is capable of spatial discrimination over a narrow line width. When the fiber under test experiences a temperature variation, it is possible to observe a red shift of the Rayleigh signature with a coefficient (10.0 pm/°C) similar to Bragg grating sensors; using a correlation method as in [53], it is then possible to obtain the temperature variation over the whole fiber length.

This principle of operation is implemented by the instrument (OBR 4600 DTS, provided by Luna Technologies) used for DTS sensing. The DTS achieves a minimum resolution of 20 µm with a maximum fiber length of 20 m and up to 1 Hz sampling rate. Performances are, however, a result
Figure 12: Photographs of the measurement chamber used for DTS experiments. (a) Measurement chamber prior to liver phantom positioning; fibers have been colored to show their positioning; (b) picture of the chamber after positioning of the liver phantom and ablation device.

of a tight trade-off between such parameters; in order to match the best RFA specifications and achieve real-time monitoring with 1s measurement rate, the spatial accuracy has been set to 200 μm. With this setting, the temperature accuracy is estimated to be 0.2 °C. The spatial resolution can be narrowed in postprocessing to 20 μm, by correlation analysis of the whole Rayleigh patterns; this feature requires extensive computations and is not available in real time. A thermistor is used to acquire the initial temperature in the tissue.

The instrument calibrated in two steps, in first place, a fiber with 20 m length and ~100% broadband reflectivity, is used to generate the reference Rayleigh signature, and subsequently the DTS is calibrated for the thermal coefficient after the fiber is positioned in the phantom.

The setup for DTS measurement, as shown in Figure 12, exploits the long-range sensing for monitoring RFA along several ablation axes. The ablation device (4 mm diameter, 1 cm electrode, as in Figures 3(a) and 9) is inserted from the bottom side of a cylindrical measurement chamber having 72 mm diameter. One single-mode fiber (Corning SMF-28) is connected to the DTS and performs temperature sensing. The fiber intersects the chamber along multiple axes, lying on the ablation plane, as in Figure 12(a); this setup allows measuring the temperature along 8 radii. Liver phantom is then positioned on top of the ablation plane, as in Figure 12(b). The measurement principle of this setup is shown in Figure 13: the DTS records the temperature profile over the whole fiber length, and the only portion including the RFA monitoring is processed to estimate the temperature variation.

Figure 14 reports the thermal map observed for each radius measured from the ablation tip, for a RFA procedure with 15 W electrical power having 269 s duration. The three-dimensional chart shows an evident similarity between the different directions; only during the liquid-to-vapor transition, which is the least predictable part of RFA [45], a perceivable difference between the different directions can be observed. Temperature recorded by the DTS system has a similar pattern with respect to thermal maps recorded with FBG and LCFBG sensors (Figures 5 and 10); in particular, temperature exhibits a quadratic increase during the first part of RFA, until reaching a plateau around 90–95 °C, and then rapidly grows before the RF generator discontinues power supply.

In order to better evaluate the shape of temperature distribution, Figure 15 reports the temperature pattern as a function of distance from the ablation device. Temperature shows an initial rapid growth, followed by a slower increase that runs through the majority of the RFA duration. During this phase, the temperature distribution follows a constant pattern, compatible with [15, 45]. As temperature approaches

Figure 13: Principle of operation of DTS measurement. The chart shows data recorded during a RFA experiment. The upper chart shows the Rayleigh signature of the whole fiber span used in the measurement. The DTS connector at 0 m length is visible, as well as the not cleaved output connector at 3.75 m distance from the DTS output. The red part of the chart contains the Rayleigh backscattering acquired in the fiber portion including the parts of fiber enclosed in the measurement chamber as in Figure 12. This part of the trace is processed by the DTS to estimate the temperature variation (bottom chart); the 8 measurement radii are visible as 4 thermal peaks.
Figure 14: Evolution of a RFA treatment, with 15 W RF power, over 8 different radii. For each radius monitored with the DTS, the temperature map is reported as a function of the distance from the ablation tip and elapsed time.

the final plateau preceding the RF power suspension, it is possible to observe a change of shape in the spatial distribution, as the heat field appears to increase its depth of penetration in the tissue, particularly for the 7–9 mm interval.

In order to evaluate the effect of the “average” RFA, Figure 16 reports the instantaneous temperature, average over 4 ablations performed with the setup as in Figure 12 on different porcine phantom samples and with the same RF power, as a function of elapsed time; the temporal axis is normalized in order to average the different ablations. The chart shows, similar to Section 3.1, that temperature rise has a different slope as a function of distance from the ablation device tip. It is also possible to notice that the ablation peak occurs with increasing delay moving away from the ablation point.

The setup reported in Figures 12–16 exploits the capability of the OFDR system for distributed sensing; this setup is
a powerful tool for multi-axes measurement. In order to validate the possibility to use the DTS in vivo, a simple measurement setup that mimics Figure 3(c) has been used, using the same ablation device as in Figures 12–16 and 15 W power and having the sensing fiber externally inserted on the ablation plane along the longitudinal axis, as in Figure 3(c). Temperature has also been recorded with two thermistors (B57861 Epcos, Munich, Germany; accuracy ± 0.2°C), one positioned internally to the hollow ablation device and one positioned externally at 3 cm distance (Figure 17).

The result shows that the DTS provides a different response to thermal variations compared to the electrical thermistor [15, 22, 27]. This is mainly due to the positioning of the thermistor inside the ablation device and the absorption of irradiance on the active sensing part [26, 27]. Such effect is partially mitigated by the higher active volume. The DTS provides a response similar to Figure 16, and it exhibits a lower response time. Measurement shows that the ablation outcome results in a small volume being treated: while the 5 mm curve clearly shows the temperature exceeding the 60°C threshold, the 10 mm curve achieves a maximum temperature of 51°C in correspondence to the ablation peak.

5. Fabry-Perot Interferometry

Pressure Sensing

5.1. Significance and Sensing Challenges. The measurement of pressure in RFA is an asset for the advanced treatment of encapsulated tumors, such as hepatocellular carcinoma (HCC) [13–16, 55], which are tumors enclosed by fibrotic connective tissues, forming a cavity between the tumor tissue and its enclosing capsule. When treated via RFA, the ablation of encapsulated tumors shows a more asymmetrical pattern [45, 56], and the ablation procedure is less effective than in tissues having a more uniform depth of penetration. Pressure is a key metric for advancing RFA in encapsulated tumors.

Tanaka et al. [56] have first introduced the intratumor pressure measurement as a parameter for RFA. However, the pressure measurement in thermal ablation, particularly RF-based, has encountered significant technological challenges mainly due to sensors characteristics.

Kotoh et al. [29] developed a setup for the intratumor pressure estimation, recording pressure values higher than 750 kPa. A microelectromechanical sensor (MEMS) has been used for pressure recording; however, the MEMS has been positioned at 3 cm distance from the ablation tip. This setup succeeds in providing a temperature-independent measurement, as the sensor is positioned outside of the area subjected to ablation, but it does not provide pressure readout at the point of treatment. Nakamuta et al. [57] have improved the measurement methodology using a phantom that models encapsulated HCC and demonstrated pressure measurement in RFA using a setup similar to [29] with a multitile ablation.
device [58]. References [29, 57] show the possibility to advance the RFA treatment, in a multistep procedure based on pressure recording; such approach is not performed in vivo, as pressure recording is not implemented at the point of treatment and MEMS sensors are not suitable for this application.

Pressure sensing in RFA monitoring is an overwhelming challenge, which requires significant advancement in sensor technologies. In particular, pressure probes need to address five specifications:

1. Both the active and passive part of the sensor, including its cables, need to have a miniature footprint, in order not to alter the ablation pattern. Sensor/cable diameters in excess of 0.25 mm can significantly change the profile of heat irradiance, either when made with dielectric material (e.g., glass fibers, which can electrically insulate the surrounding tissue) or conductive material (e.g., metal packaging of sensors [28], which can bias the irradiance in the cable direction altering the symmetry of the ablation pattern).

2. The sensor has to be biocompatible for in vivo use; typical reference standard is ISO 10993 [59].

3. The sensor acts in close proximity to the RF ablation tip; hence, MRI compatibility needs to be guaranteed. The use of a small sensor is also advantageous as a small active volume results in a lower biasing induced by RF irradiance on the sensor tip.

4. The sensor needs to be thermally insensitive. With most sensing technologies, including MEMS and FOS, pressure probes are cross-sensitive to temperature; in RFA, both temperature and pressure exhibit steep variations throughout the duration of the measurement; hence, thermal insensitivity needs to be guaranteed. On the other side, temperature gradients occurring at the point of RFA treatment, as documented in Sections 3 and 4, make it extremely arduous to develop a thermal compensation technique.

5. The probe needs to have mechanical strength sufficient to sustain the penetration into the tissue and the extraction upon completion of RFA.

MEMS pressure sensors, as [29, 57], are typically MRI compatible and biocompatible, but they are temperature compensated only up to ~50°C and their footprint has excessive size compared to the volume of tissue exposed to RFA. Fiber-optic sensors [25] represent a more valuable alternative for this task.

Extrinsic Fabry-Perot interferometry (EFPI) [18, 30, 60–65] is the most popular principle of operation for fiber-optic pressure sensing. EFPI sensors are substantially designed as a Fabry-Perot cavity on the tip of an optical fiber, sealed with a pressure-sensing diaphragm [30, 60–66]. By using optical fibers, MRI compatibility is guaranteed [30, 59], and the size of the sensor is suitable for RFA. On the other side, the vast majority of the solutions proposed for EFPI sensors fail to address altogether the other challenges for RFA sensing.

In the first place, sensors based on epoxy, typically UV curable adhesives [63], are often not compliant to ISO 10993 standard and not resistant to high temperatures. Viable solutions include the realization of an all-glass sensor [60, 64, 66] that uses quartz glass or an optical fiber as diaphragm, metal diaphragm as in [67], which, however, may be biased by the absorption of RF radiation on the diaphragm, or an organic material layer as diaphragm [68]. Pressure sensitivity increases by narrowing the thickness of the diaphragm [61, 63, 64]; however, sub-μm thick diaphragms as [66, 67] or bendable materials [68] show excessive fragility when inserted in the tissue, even with external catheterization.

Temperature compensation has been introduced by Bremer et al. [61, 69] in EFPI sensors, using an inline FBG sensor to compensate for temperature variations from an EFPI sensor, sensitive to both pressure and temperature; this approach has been then continued in [18, 20, 63, 70] and is currently an acquaintance for EFPI sensors. The all-glass sensor proposed in [61] has a thermal detuning coefficient 1.6 kPa/°C, which is the typical value for glass EFPI probes; the epoxy-based sensor proposed by Bae and Yu [63] has a thermal cross-sensitivity one order of magnitude higher. In this framework, even if the FBG sensor can compensate the temperature variations, the length of the FBG and its displacement from the fiber tip containing the EFPI are not negligible; indeed, the temperature recorded by the FBG is not the same value at which the EFPI is exposed to.

In 2014, Tosi et al. [18] proposed an EFPI/FBG sensor fabrication that records the smallest cross-sensitivity, achieving a thermal detuning of 0.032 kPa/°C. This design is based on the fabrication of an EFPI probe with a quartz glass capillary, over a FBG sensor previously inscribed in a single-mode fiber. A novel approach, more recently introduced by Poegegel et al. [20] reverses the fabrication process, inscribing a 1 mm FBG sensor on an EFPI; this approach allows inscribing the grating of the exact tip of the probe, leaving no passive regions between the EFPI and the FBG. These approaches allow pressure sensing in RFA, as they comply with all the requirements for operation in thermal ablation.

5.2. Fabrication and Characterization. Figure 18(a) shows the fabrication of the EFPI/FBG probe [18, 61, 63, 69, 70]. In the first place, a FBG is inscribed in a standard single-mode fiber (SMF-28, 10/125 μm), using UV laser and a phase mask [32, 71]. The grating has 4 mm length, with 1540.5 nm wavelength. After cleaving the SMF fiber in proximity of the first place, a FBG is inscribed in a standard single-mode fiber (SMF-28, 10/125 μm), using UV laser and a phase mask [32, 71]. The grating has 4 mm length, with 1540.5 nm wavelength. After cleaving the SMF fiber in proximity of the grating of the exact tip of the probe, leaving no passive regions between the EFPI and the FBG. During this operation, the reflectivity of the FBG reduces from 95% to 4%. The capillary is then hand polished (λ/10) and spliced to a multimode fiber (62.5/200 μm) which serves as diaphragm; this procedure seals an air-gap Fabry-Perot cavity on the tip of the transmission fiber, with length 20–60 μm. In order to functionalize the structure for pressure sensing, the thickness of the diaphragm has to be reduced [61]; this task is accomplished by first hand polishing the diaphragm and then controlled etching in hydrofluoric acid as in [19]. All the splicing operations are performed with a manual splicer (Siecor M91).
Figure 18: Schematic of the fabrication technique for the EFPI/FBG probe. (a) Fabrication of an EFPI probe over a preinscribed FBG sensor. In the first step, the FBG is fabricated with phase-mask technique on a standard single-mode fiber [18]; then, the fiber is spliced to a hollow glass capillary, and the resulting structure is spliced to a multimode fiber serving as diaphragm. Finally, etching in hydrofluoric (HF) acid [19] reduced the diaphragm thickness for pressure sensitivity. (b) Fabrication of an EFPI probe with subsequent inscription of the FBG: in this case, the sensor is built as in (a), steps 2–4, and the FBG is inscribed on the sensor tip as in [20].

Figure 19: EFPI/FBG probe spectra. (a) Sensor fabricated with the classical method, as in Figure 18(a); (b) postinscription of FBG with the point-by-point method, as in Figure 18(b).

An alternative fabrication technique has been recently demonstrated by Poeggel et al. [20]. The EFPI sensor is designed as in Figure 18(a), obtaining a pressure probe. Then, the FBG is fabricated using a femtosecond laser and a point-by-point inscription [72]. While the previous fabrication method cannot allow a perfect positioning of the FBG on the fiber tip, as the splicing procedure performed near the air-gap tends to damage the FBG, this novel approach allows inscribing the FBG exactly on the tip of the probe. Hence, this second approach minimizes the distance between the probe tip and the FBG tail.

Figure 19 shows the spectra of the EFPI/FBG probes fabricated with both methods. The classical inscription method provides a clean spectrum, with a neat separation of the EFPI broad spectrum from the FBG narrow spectrum. The EFPI has a low finesse (0.083), given by a low 2% reflectivity. The postinscription of the FBG has the advantage of fabricating a shorter FBG on the fiber tip, more suitable for RFA; but the
resulting spectrum has a more wrinkled pattern [72]. It is still possible to discriminate the FBG from the EFPI, but pressure measurement achieves better accuracy when based on a clean spectrum.

A probe fabricated with the classical method has been used in RFA experiments; Figure 20 shows the compression of the air-gap as a function of the applied pressure (0–70 kPa range). The EFPI sensor has a sensitivity of 1.60 nm/kPa, compatible with [61, 63], and linearity coefficient 0.9997. The choice of the glass capillary, as outlined in [18], reduces the thermal sensitivity of the EFPI to 51.9 pm/°C, over 17–82°C range. This value is over 30 times smaller than that in [61] (1.7 nm/°C) and 300 times smaller than that in [63] (15.8 nm/°C). The FBG has a thermooptic coefficient of 10.7 pm/°C and negligible pressure sensitivity [63]. This sensor achieves dual pressure/temperature measurement with accuracy of 40 Pa and 0.1°C, respectively.

5.3. Experimental Results. Two experimental setups (pictures in Figure 21) have been designed for RFA experiment, following the approach used for FBG, LCFBG, and DTS sensing. In the first setup, RFA is performed with a hollow needle, 3 mm diameter and 1 cm active length, which penetrates the tissue with 2 cm approximate depth. The EFPI/FBG probe is inserted through the RFA device, and the fiber tip is located at approximately 1 mm distance from the tip of the active electrode. This setup mimics Figures 3(a)-3(b) and 9(a). The second setup, which is closer to Figures 3(c), 9(b), and 12, encloses the porcine liver tissue in an XPS chamber covered with a plexiglass layer, as in Figure 3(c). The EFPI/FBG probe is inserted from the side, orthogonally to the RFA needle. The picture in Figure 21 shows the pattern of ablation after the procedure is completed. Two experiments have been performed, having the fiber tip positioned at 5 mm and 8 mm from the tip of the electrode. In order to better visualize the
pressure pattern, these RFA experiments have been carried out with an inferior RF power (5 W) to extend the duration of ablation treatments.

The result of a RFA procedure carried out with the first experimental setup is shown in Figure 22, whereas both pressure and temperature traces are displayed. After the initial setup, temperature rises quadratically (as in previous experiments), while pressure appears to be floating, initially reaching negative values (−13 kPa) and then achieving a peak at 39 kPa. As temperature rises to ~80 ºC, pressure continues its oscillation reaching a maximum value close to 50 kPa. At this stage, temperature starts a rapid growth reaching a maximum value of 164 ºC; the pressure trace, after an initial decrease to 8 kPa, rapidly increases first to 125 kPa and then to 162 kPa, 3 s after the temperature peak is recorded.

This chart is significantly different from the previous measurement by Kotoh et al. [29, 57]. Quantitatively, the value of 750 kPa recorded at 3 cm distance from the ablation tip has not been recorded, and maximum pressure recorded is 162 kPa (1.62 bar). The trace also has a different profile: while the fluctuations can be partially explained by the change of refractive index of the medium surrounding the EFPI probe, pressure exhibits two almost instantaneous rise, the second one being recorded with a delay with respect to the temperature peak.

Figure 23 reports, for the experiment in Figure 21, the pressure values as a function of temperature (PT chart). In this plot, it is possible to distinguish how the two quantities are correlated. In the initial phase (1), temperature rises from ambient to 101 ºC, while pressure fluctuates from the initial zero to negative and positive values, −13 kPa to 50 kPa. Then (2), the chart experiences an almost vertical transition: pressure rapidly grows to 92 kPa within few seconds, in which temperature rises at a slower pace. In the region (3), both pressure and temperature exhibit a rapid change, in correspondence to the liquid-to-vapor occurring in the liver tissue: temperature quickly rises to its maximum value, while pressure first increases to 125 kPa and then decreases to 88 kPa. The last two movements on the PT chart occur at a very rapid pace: temperature drops with an instantaneous rate, corresponding to a horizontal movement (4), followed by a vertical shift (5) in correspondence to the pressure peak, occurring when the liver temperature is already <40 ºC.

The results of three RFA experiments are reported in Figure 24 reports, for the experiment with setups as in Figure 21. In first place, it is possible to notice that the pressure values are correlated with the distance from the fiber tip, as for temperature variations; maximum pressure values of 162 kPa, 33 kPa, and 6 kPa have been recorded at 1 mm, 5 mm, and 8 mm distance from the fiber tip to the electrode tip (31 ºC, 46 ºC, and 164 ºC maximum temperature, resp.). In both the first and second experiment, it is also possible to notice that pressure exhibits a similar trend: after an initial stabilization, with an average rise, pressure increases in correspondence to the phase transition of the liver tissue, with a slight delay with respect to temperature rise. After temperature reaches its peak, pressure continues rising and the maximum value is reached with some delay with respect to the temperature maximum. The pressure peak has been recorded for normalized time equal to 1.05 (first experiment) and 1.07 (second experiment). At 8 mm distance, the pressure and temperature variations are significantly smaller.

The PT chart for the RFA experiments carried out with the setup as in Figure 21(b) and with the EFPI/FBG probe at a further distance from the ablation tip shows a different pattern. Figure 25 shows the PT chart for the second RFA experiment in Figure 24. In the first place, it is possible to notice that the pattern is more linear, and the amplitude of pressure fluctuations is lower; this can be partially explained by the lower variations of refractive index (RI) surrounding the probe but also to the better enclosing of the liver that prevents pressure from rising and decaying. Compared to Figure 23, most of the trace appears to be confined in region (1), whereas both temperature and pressure rise at a steady rate. When temperature reaches the 40–44 ºC interval, pressure appears to rise linearly reaching 23 kPa value. As temperature starts to decrease, we observe a pressure rise corresponding to a vertical transition on the PT chart; this is analogous to region (5) in Figure 25, that is, a rapid pressure rise, when temperature is reducing. Qualitatively, the chart
appears to have fewer transitions and is missing regions (1), (2), and (3).

6. Discussion

6.1. Technology Review and Developments. The optical fiber sensor technologies used in this work provide an all-encompassing view of the possible strategies for monitoring of biophysical parameters in RFA procedures. Each approach provides different performance levels and can be dedicated to a specific task in RFA monitoring. Table 1 reviews the performance of all sensors.

FBG sensors provide the easiest way for the monitoring of a small number of sensing points, using inline sensing. FBGs have typical length of 1–4 mm, which is not negligible considering the thermal gradients generated in RFA [32]. Using standard FBGs [35], the fabrication of a sensor array is limited by splicing or other fabrication techniques to ∼2 cm spacing between each FBG. Dense FBG sensors arrays have been proposed as in [27], achieving up to 3 sensors/cm; this fabrication cost for such structures is however excessive for thermal ablation. The fabrication of FBGs on a drawing tower (DTGs) is a flexible alternative, which can have potential in RFA [26, 37]. Most DTG fabrication facilities are limited to 5 mm minimum length and 10 mm minimum sensor spacing, for the fabrication of low-cost sensors; unlike traditional FBGs, the fabrication cost of a DTG is scalable with the number of sensors, as the fabrication of FBG arrays has a high fixed cost and a marginal variable cost depending on the number of sensors. The main asset of grating sensors relies on the consolidated techniques for interrogation of FBG arrays, which is nowadays implemented in a broad number of commercial devices.

At the present state of the art, the poor resolution of FBG/DTG sensors is a key limitation in thermal ablation. However, this scenario is likely to change within the next
few years, with the advent of point-by-point FBG inscription [72] that can potentially fabricate grating structures with a narrow resolution and with potential improvements to the DTG fabrication approach [36, 37]. Currently, DTGs are also applied for all-grating fibers [73, 74], which extend FBG sensing to an entire fiber span with distributed integration but are still limited in resolution by the spatial resolution of FBGs. Most DTG fabrications are already established for fibers with 80 \(\mu\)m diameter, and 40 \(\mu\)m fibers are already commercially available and soon utilisable in FBG sensors, reducing the active volume of each sensing point.

The LCFBG approach, on the other hand, shows a better potential for distributed sensing [17]. It makes use of the same optical hardware as FBG sensors, using the same interrogators, but achieves a sub-0.1 mm theoretical resolution. LCFBG sensors are more expensive than DTG arrays, but they use similar fabrication techniques and can benefit as well from potential improvements of point-by-point inscription; in addition, chirped gratings are commonly employed in the fiber-optic communication industry as broadband reflectors or gain-flattening filters, and therefore it is likely that, once the technology is established, their manufacturing cost can significantly reduce. The FBG used in experiments has 1.5 cm length, but it is possible to fabricate LCFBGs having length up to 5 cm and ~80 nm bandwidth, still achieving ~0.1 mm spatial resolution.

The main drawback of LCFBGs is that, currently, there is no reliable technique for their detection in the spectrum domain. Approaches like [42, 44] require an overwhelming complexity and are not affordable in real-time operation. Simplified approaches substantially turn the LCFBG into a few-point sensor [41], which does not offer significant advantages over FBG sensors. In particular, the main problem of all the proposed techniques, including the simplified approach proposed in this work, is that the entire spectrum needs to be detected as a whole, as the temperature estimated in each sensing point depends on the values recording in all the adjacent points.

This statement is well demonstrated in Figure 26, which compares different temperature readings performed with LCFBG, DTS, and DTG-FBG array. Although the comparison is only for qualitative purpose, as it includes different RFA experiments on several porcine phantoms and measurements are recorded along different axes, it shows that the LCFBG has a different qualitative behavior with respect to the other technologies, which however show similar readings. The simplified approach proposed in [17] for distributed temperature estimation needs to be improved in order to make the LCFBG a reliable RFA-prone sensor. However, once such technique would be established, the sensing approach based on chirped gratings may have significant advantages over DTS: its performances (measurement rate, accuracy, and resolution) are not based on a trade-off but can be individually selected and improved by changing the optical hardware. In addition, the LCFBG sensor is not significantly affected by poor cleaving at the fiber end, which is a weakness of DTS.

The DTS, on the other side, gains the benefits of allowing distributed sensing over long fiber spans. Currently, OFDR technologies allow sensing over up to 20–70 m length and are often used for characterizing optical connectors [52]. The approach proposed by Gifford et al. [53] allows temperature sensing with 20 \(\mu\)m (offline)–200 \(\mu\)m (online) spatial resolution. The main attractiveness of the DTS is that, although the instrument is expensive, having a cost that usually overcomes the RF ablation equipment, the sensor is instead a standard optical fiber, having the lowest cost among any fiber-optic technologies. Although performances depend on the tight trade-off between the different parameters, the settings used in experiments are sufficient for general-purpose RFA monitoring.

DTS is an excellent instrument for ex vivo monitoring and can be used for the realization of complex setups as Figure 12 which can really advance the state of the art of RFA modeling.
such as the European FP7 project IMPPACT [22]. The main drawback of the DTS system is that the backreflection occurring at the fiber end substantially overlaps the trace at the fiber tip (as is clearly visible in Figure 13), which is the most significant sensing information as this part of the sensor is usually located in correspondence to the ablation tip. Although a proper fiber cleave can solve this problem, when the fiber penetrates the tissue its surface gets microdamages, neutralizing the initial cleaving quality. A possible solution is the introduction of the fiber in a microcatheter, which sustains the penetration while preserving the cleaving quality.

The three technologies presented in this work require a different calibration procedure. FBGs are linear sensors that require the characterization of the thermal coefficient, which is usually carried out prior to their use and depends almost entirely on the glass compound [33]. It is possible to obtain then the absolute temperature measurement when the sensor is initialized, either using an external sensor (ex vivo) or using the temperature reference value 36.8 ± 0.4°C (in vivo). The calibration of a LCFBG requires the determination of the thermal coefficient and the temperature offset, as the FBG, but it also needs the characterization of the LCFBG spectrum in uniform heating conditions, that is, when the whole LCFBG is exposed to the same local temperature [17, Figure 7]. This characterization can be performed in a temperature chamber but may substantially differ from each LCFBG sensor as it can depend on spectral ripples, chirp coefficient, and other fabrication parameters. The DTS requires the acquisition of the reference chaotic Rayleigh signature, which is then used to estimate the wavelength shift corresponding to temperature variations in each measurement point [53]. This task is performed by calibrating the whole instrument in reference conditions, whereas the whole temperature distribution is known (e.g., constant temperature through all the fiber). However, the calibration of the instrument prior to fiber insertion is arduous, since fiber bends and eventual backreflections often occurring with the insertion of the fiber at the point of treatment may substantially detune the measurement. A simple, yet effective, calibration technique is to impose the temperature distribution in a specific portion of the fiber link, which is assumed not to have optical losses; this can be a viable solution for the application of the DTS in vivo, assuming that the temperature of the fiber is constant at the reference value 36.8°C and neglecting the other parts of the fiber span.

As outlined in Section 5.1, the state of the art of pressure sensor does not offer valuable alternatives to the sensor presented in this work; to the best of our knowledge, no other solution can offer thermal insensitiveness, biocompatibility, miniature size, and tensile strength, which are all essential requirements for RFA. The pressure measurements reported have more significance than [29], as they are performed at the point of treatment rather than at distance. The future work will be addressed to improving the fabrication of sensors to high-volume manufacturing rates, providing a better insulation of the fiber tip from the surrounding refractive index to avoid pressure fluctuations and advancing the clinical knowledge of advanced treatment of encapsulated tumors supported by fiber-optic probes. EFPI/FBG integrated probes offer the advantage of integrating a temperature sensor and give access to the PT chart (Figures 23 and 25). EFPI/FBG probes make use of the same interrogation setup of FBG and LCFBG sensors, and, with the technique proposed in [75], the pressure accuracy is as low as 6 Pa.

All measurements carried out in experimental setups are based on single-channel detection; however, interrogation of 4–16 channels with optical switching can extend the sensing capabilities. Using the most recent fiber-optic switches with 20 ms switching time and building custom-made software for interrogation using off-the-shelf hardware, it is possible to obtain 1 Hz measurement rate with up to 16 channels. Such feature is not available for DTS systems, in which the measurement duration is tied to the accuracy/resolution performance levels.

6.2. Smart Ablation. The ultimate goal of the application of fiber-optic sensors in RFA is to gain the capability of performing ablation procedures based on sensors readout, rather than imaging, and dynamically change the parameters of ablation in a closed-loop way. Such technique, hereby called "smart ablation," is currently limited by the lack of sensing technologies; this work aims at filling the gap, highlighting sensing technologies that are compatible with the application in RFA. A possible implementation of the smart ablation concept is sketched in Figure 27.

Sensors can be (1) installed on each tip of a multitet device [21, 57, 58], (2) inserted through a hollow ablation device in an electrically passive region, or (3) externally inserted with a catheter, which can eventually be removed after insertion in vivo. The latter option has been experimentally tested, with positive results [18], showing that the fiber can always be extracted from the tissue even after the catheter has been removed. A combination of LCFBG or FBG array sensors with EFPI sensors can be used. All these sensor typologies perform wavelength-selective operations and, therefore, can be all interrogated with a white-light system (WLS) operating in the near infrared, as in Figures 2 and 7, which is based on a broadband source, a coupler, and a spectrometer for spectral detection. A single WLS system can be interrogated up to 4–8 channels, each including a combination of sensors. DTS can provide an alternative technique for temperature or strain detection, operating
with high density over a single fiber. Combining WLS-based sensors and DTS, it is possible to record up to several hundreds of sensing data at the point of treatment, applying the lab-on-a-fiber concept with high density.

Using the sensors data, it is then possible to design a data fusion system that collects the biophysical data and uses them to adjust the parameters of RFA. The data fusion system can also incorporate imaging [23, 24] and modeling [22]. Haemmerich and Schutt [10] demonstrated that the ablation pattern has a dependence on the RF frequency, which suggests that smart ablation can be effective for advanced treatment of tumors. Future research will also focus on modifying the ablation device, enabling the possibility to change its position and shape dynamically through RFA, following the robotic microsurgery concepts [76, 77]. The smart ablation concept may also be extended to microwave ablation [78, 79] and cryoablation [80] of tumors.

7. Conclusions

The present work presents and reviews the use of optical fiber sensors in cancer thermotherapy. The specific medical scenario discussed in the present work involves the thermal ablation of liver tumors with a radiofrequency source. RFA procedure makes use of a percutaneous ablation device and hence can accept only miniature sensors such as optical fiber sensors as monitoring tool; on the other side, RFA is vulnerable, for the treatment of mid-to-large tumors, to the change of electrical impedance of the target tissue when temperatures around ~100°C are reached in proximity of the ablation device.

The present sensor monitors biophysical parameters: temperature distribution and pressure. Temperature distribution, sampled with the narrowest possible spatial resolution, is a key indicator of the mortality rate of cells in every portion of the tumor. Three technologies are presented: FBG arrays achieve a 1 cm resolution but allow straightforward measurement and interpretation; a LCFCBG sensor on the other hand achieves 75 μm resolution, but it is highly dependent on the interpretation of the optical spectrum, which is suboptimal with the algorithms presented to date; high-density DTS based on OFDR allows achieving resolution of 20–200 μm, with a tight performance trade-off (accuracy, resolution, and speed), but its performances ex vivo are hardly replicable in vivo unless special catheterizations are applied. Pressure is another key biophysical parameter, mainly for the optimal treatment of encapsulated tumors such as HCC. The authors have developed an EFPI pressure sensor, integrated with an FBG temperature sensor, which records, to the best of our knowledge, the lowest cross-sensitivity parameter. This achievement enables pressure sensing in RFA at the point of treatment, and preliminary experiments show a pressure rise in RFA procedures which depends on the distance from the ablation tip. This result is the first step towards adjusting RFA in encapsulated tumors, such as HCCs, to pressure readout.

In conclusion, optical fiber sensors can substantially advance the treatment of tumors via thermal ablation, providing real-time distributed or quasidistributed biophysical sensing at the point of treatment. The collection of such big data can be exploited to design a smart ablation system, whereas sensor readouts are used to dynamically adjust the pattern of ablation in real-time.

Conflict of Interests

The authors declare that there is no conflict of interests regarding the publication of this paper.

Acknowledgments

The authors acknowledge the contribution of Sven Poege, Gabriel Leen, and Elfed Lewis (University of Limerick, Optical Fibre Sensors Research Centre); Mario Gallati and Giovanni Braschi (Università di Pavia, Dipartimento di Ingegneria Civile ed Architettura); Giorgio Busca (Politecnico di Milano, Dipartimento di Meccanica); and Sandro Rossi (Policlinico San Matteo). The authors acknowledge Ian Shannan and Maurizio Chiani (Luna Technologies) for providing a DTS unit and Hugo Mertens, Johan Vlekken, and Eric Lindner (FBGS International) for supplying ormoceramic fibers. Research has been funded by Marie Curie actions (MC-IF-299985), Science Foundation Ireland (10/RFP/ECE2898), Enterprise Ireland (CF-2014-0107-Y), and Fondazione per la Cura Mini-Invasiva dei Tumori.

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