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Communication

On the Use of Impedance Detuning for Gastrointestinal Segment Tracking of Ingestible Capsules

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Abstract—During their travel through the gastrointestinal tract, ingestible antennas encounter detuning in their impedance response due to varying electromagnetic properties of the surrounding tissues. This paper investigates the possibility of using this impedance detuning to detect in which segment of the gastrointestinal tract – stomach, small intestine, or large intestine – the capsule is located. Meandered dipole antennas operating in the 433 MHz Industrial, Scientific, and Medical Band are designed for this purpose. The antennas conform to the inner surface of 3D-printed polylactic-acid capsules with a shell thickness of 0.6 or 0.4 mm. The impedance response is first optimized numerically in a homogeneous cylindrical phantom with time-averaged electromagnetic properties. The magnitude and the phase of the reflection coefficient are then obtained in different tissues and compared with simulations and measurements. The experimental demonstration is carried out first using tissue-mimicking liquids and then in a recently deceased *ex vivo* porcine model. The minimum change in the phase between different gastrointestinal tissues was determined to be around 10° in the porcine model, indicating that the changes in the impedance response, particularly the changes in the phase, provide sufficient information to follow the position of the capsule in the gastrointestinal tract.

Index Terms—Conformal antennas, dipole antennas, impedance detuning, in-body, ingestible devices, *in vivo* applications.

I. INTRODUCTION

In recent years, ingestible devices have been widely used in diverse medical applications, such as monitoring of physiological data and wireless endoscopy [1]–[3]. Due to the variety of possible applications and compact structure of these devices, the ingestible bioelectronics have the potential to substantially improve the diagnostics and therapies of the gastrointestinal (GI) tract [4].

Ingestible devices consist of several components integrated into the same capsule [5]. The antenna is one of the key components as the quality of the communication link with external devices strongly depends on its performance [6], [7]. Therefore, different types of antennas, such as patch [8]–[10], spiral [11], helical [12], [13], and differentially-fed antennas [14], [15] have been implemented in ingestible capsules. Note

that the ingestible antenna design is a challenging task due to the variety of loss mechanisms encountered in their operation. These losses can be classified as near-field, propagation, reflection, and detuning losses [16]–[19]. The latest corresponds to the possible changes in the impedance response of the antenna caused by the physical and electromagnetic (EM) variations in the surrounding biological environment as the capsule travels through the GI tract. Several studies in the literature examined these changes at different frequencies, such as 403.5 MHz [20], 433 MHz [21], [22], and 2.45 GHz [23], [24]. Moreover, various robust ingestible antenna designs were proposed to mitigate this loss component [25]–[32].

Recent technological breakthroughs in terms of miniaturization, artificial intelligence, and power saving have enabled innovative companies to develop ingestible systems for monitoring physiological data [33], [34]. The next generation of ingestible bioelectronic systems aims to provide information beyond this intent by performing accurate measurements all along the GI tract. For instance, the real-time localization of a capsule inside the GI tract can be a useful and innovative measurement to (i) optimize a device to investigate the middle of the track which is not accessible with probes (feeding tube, rectal probe), (ii) correlate a measurement and a position to determine the segment duration in order to identify specific pathologies such as gastroparesis, (iii) deliver efficiently an active substance at the right position. The smart capsules designed within this context can offer an ergonomic and accurate medical system.

In this context, this paper investigates the possibility of using the impedance detuning caused by the varying EM properties of the GI tract to distinguish the GI tissues (i.e. stomach, small intestine, or large intestine) for localization purposes. For this investigation, impedance responses (both magnitude and phase of the reflection coefficient) of capsule-integrated meandered dipole antennas in different GI tissues are compared with simulations in numerical phantoms representing different GI tissues and with measurements in tissue-mimicking liquids as well as in an *ex vivo* porcine model. To the authors' best knowledge, it is the first time in the literature where impedance responses of ingestible antennas are investigated for the purpose of GI segment tracking.

This communication is organized as follows: First, Section II describes the antenna and simulation models, and presents the simulation results. Then, Section III explains the prototyping process, and discusses the measurement results. Finally, the paper concludes in Section IV.

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II. NUMERICAL STUDY

A. Antenna Design and Modeling

A capsule-integrated ingestible antenna is designed for the intended purpose. A meandered dipole antenna is preferred for the design since its radiation mechanisms and the near field behavior have been widely studied and better understood than the complex structures [16], [35]. The designed antenna, as visualized in Fig. 1a, is a meandered dipole antenna operating in the 433 MHz Industrial, Scientific, and Medical (ISM) band. It is designed on a 100- μm -thick Rogers CLTE-MW substrate ($\epsilon_r = 2.97$) [36], which conforms to the inner surface of the polylactic-acid (PLA, $\epsilon_r = 2.7$, $\delta = 0.003$) capsule as shown in Fig. 1b. The capsule is also filled with a PLA cylinder to fix the antenna in its cylindrical shape and has two thicknesses: $t = 0.6$ mm or $t = 0.4$ mm. It is placed in the middle of a cylindrical homogeneous phantom, as seen in Fig. 1c. The trace length is numerically optimized for the two shell thicknesses in the phantom having time-averaged EM properties of the GI tract ($\epsilon_r = 63$, $\sigma = 1.02$ S/m). The detailed description of the time-averaged phantom can be found in [24], [37]. The optimized values for the trace length are 16.4 mm for $t = 0.6$ mm and 14 mm for $t = 0.4$ mm. The values of the other parameters are the same for two shell thicknesses and are shown in Fig. 1a. The parameter N indicates the number of turns as explained in [38] and is equal to 7 for both models. Note that an offset feed is required to better match the antennas to 50 Ω due to long antenna meandering [39].

B. Numerical Results

After optimizing the antennas, the possibility of using the impedance detuning is examined by simulating the antennas in numerical phantoms representing 3 tissues in the GI tract: stomach, small intestine, and large intestine. The EM properties of these tissues at 434 MHz are tabulated in Table I [40]. The simulated magnitude and phase of the reflection coefficient are shown in Fig 2. From the results, it can be seen that the minimum difference in the magnitude observed between 3 GI tissues at 434 MHz is 0.9 dB for $t = 0.6$ mm and 1.1 dB for $t = 0.4$ mm. As for the phase, it is 16.9° for $t = 0.6$ mm and 19.7° for $t = 0.4$ mm. As these values are sufficiently large to distinguish the tissues, it can be stated that the change in the impedance response of the antenna in different GI tissues can be utilized to determine the segment in which the capsule is located. Note that it is more convenient to track the change in the phase for the intended purpose, as it is more dominant than the change in the magnitude.

III. PROTOTYPING AND MEASUREMENTS

The measurements were carried out using two different measurement setups. In the first setup, tissue-mimicking liquids were used. In the second setup, *ex vivo* measurements were performed using a porcine model.

A. Prototyping

The antennas and 3D-printed capsules were fabricated for the measurements, as shown in Fig. 3a and Fig. 3b, respectively. The antennas were soldered to the coaxial cable and

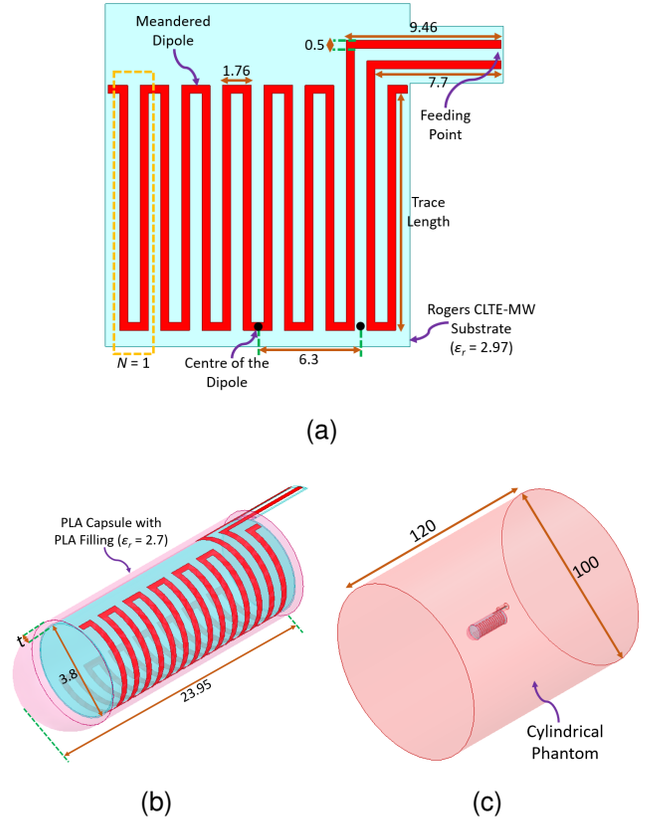


Fig. 1. Model of the antenna, the capsule and the numerical setup (units: mm). (a) Meandered dipole antenna model. (b) Antenna conforming to the inner surface of the PLA capsule. (c) Capsule placed inside the cylindrical numerical phantom for simulations.

TABLE I
EM PROPERTIES OF THE GI TISSUES AT 434 MHz AND THE AMOUNT OF EACH INGREDIENT (IN GRAMS) USED PER 100 G MIXTURE FOR THE FABRICATION OF THE TISSUE-MIMICKING LIQUIDS

	Average	Stomach	Small Intestine	Large Intestine
ϵ_r	63	67.2	65.3	62
σ (S/m)	1.02	1.01	1.92	0.87
Water	53.96	59.13	54.59	52.97
Sugar	44.6	39.7	42.6	45.8
Salt	1.44	1.17	2.81	1.23

placed inside the capsules. The side of the capsules was covered with Araldite 2012 epoxy resin to make a watertight seal. As mentioned previously, the main purpose of this study is to compare the impedance responses of the antennas in different GI tissues. However, with the introduction of the coaxial cable, the EM changes in the environment around the cable also affect the impedance responses, decreasing the accuracy of the comparison. Therefore, ferrite rings, which help eliminate this undesired effect by attenuating the magnetic fields created between the cable and the antenna, were placed at the antenna end of the cable. Note that the cable affects the operation of the antennas even with the use of ferrite rings. However, the ferrite rings stabilize this effect in different tissues by making it independent of the surrounding environment, which is sufficient for the aim of this work. The complete prototype

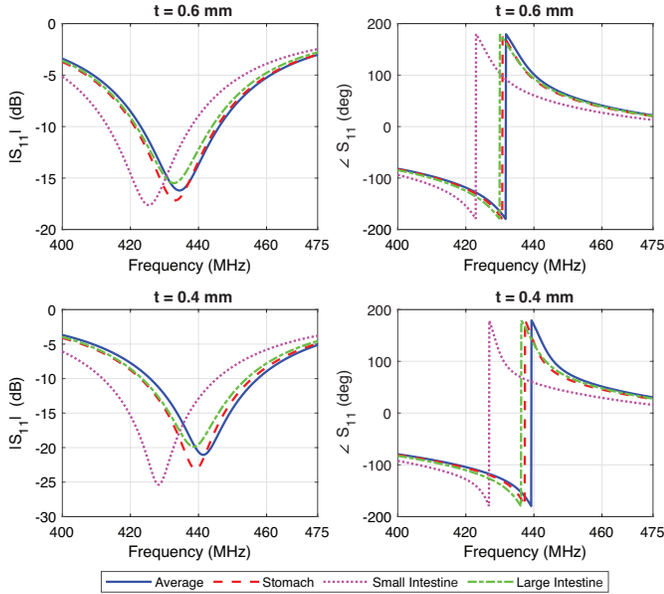


Fig. 2. Simulation results for the magnitude and the phase of the reflection coefficient obtained in numerical phantoms representing different GI tissues for two shell thicknesses.

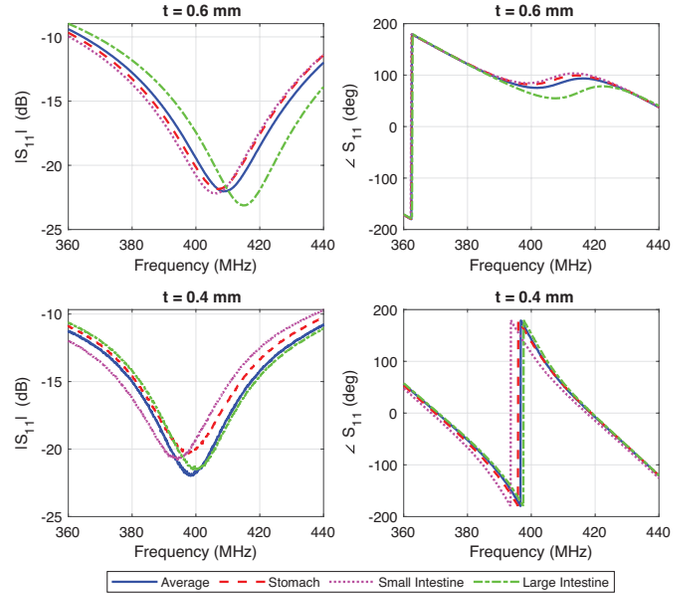


Fig. 4. Measurement results for the magnitude and the phase of the reflection coefficient obtained in tissue-mimicking liquids for two shell thicknesses.

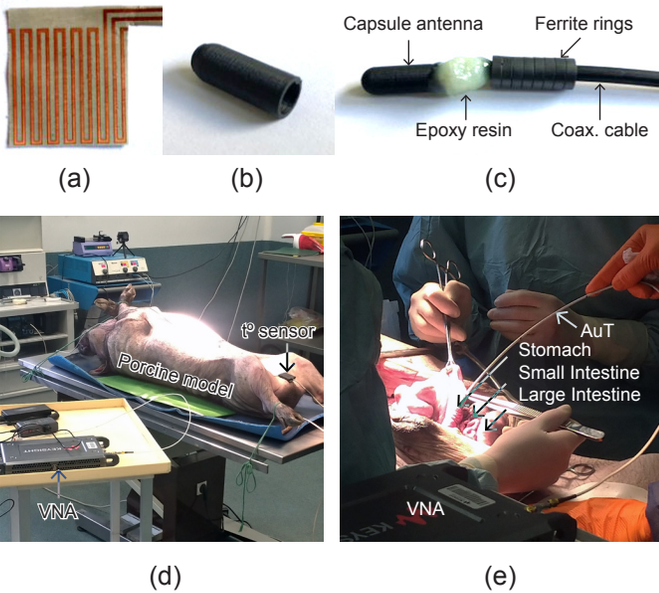


Fig. 3. Different parts of the prototype and the porcine model. (a) Printed antenna. (b) 3D-printed PLA capsule. (c) Final antenna under test (AuT) with the ferrite rings. (d) General view of porcine model used in the *ex vivo* measurements. (e) AuT placed inside the porcine model.

with the ferrite rings can be seen in Fig. 3c.

B. Measurements in Tissue-Mimicking Liquids

For the first measurement setup, tissue-mimicking liquids were fabricated with deionized water, sugar, and salt using the method proposed in [27]. The amount of each ingredient used for 100 g tissues is tabulated in Table I. The EM properties of the fabricated liquids were validated at 434 MHz with a SPEAG DAK-12 probe [41]. Fig. 4 shows the measurement results in the prepared liquids. It can be observed that the

operating frequency in the liquid with time-averaged EM properties is shifted to 408.6 MHz for $t = 0.6$ mm and 398.3 MHz for $t = 0.4$ mm due to the effect of the cable. At these resonant frequencies, the minimum difference in the phase observed between 3 GI tissues is 5.3° for $t = 0.6$ mm and 16.3° for $t = 0.4$ mm. Similar to the simulation results, these values are sufficiently large to identify the tissue in which the capsule is located, supporting the idea presented in this work.

C. Ex vivo Measurements

In the second measurement setup, *ex vivo* measurements were performed using the porcine model shown in Fig. 3d. Experimental validation of the proposed EM-sensing with ingestible capsules requires measurements using recently deceased tissues as the EM properties might quickly evolve post-mortem [42]. Thus, *Ex vivo* measurements on a 50-kg live body-weight Yucatan miniature pig were performed at the experimental facilities of the Unité Expérimentale Physiologie et Phénotypage des Porcs in Saint-Gilles, France [43].

After premedication with ketamine (15 mg/kg IM), a 50-kg live body-weight Yucatan miniature pig was euthanized by intravenous injection of 5 ml of T61 – a nonbarbiturate, non-narcotic combination consisting of 3 compounds: embutamide (anesthetic), mebenzonium iodide (paralytic), and tetracaine hydrochloride (potent local anesthetic). Once euthanized, the animal was placed in dorsal recumbency. An incision (about 30 cm) was made in the lower abdominal wall ventral median following the linea alba starting 100 mm behind the breastbone, opening the abdominal cavity. The stomach (M1), the apical part of the duodenum (M2), the terminal part of the ileum (at the beginning: M3 and at the end: M4), and the terminal part of the colon (M5) were located and exteriorized. At each location, 1 cm incisions were made to insert the prototypes

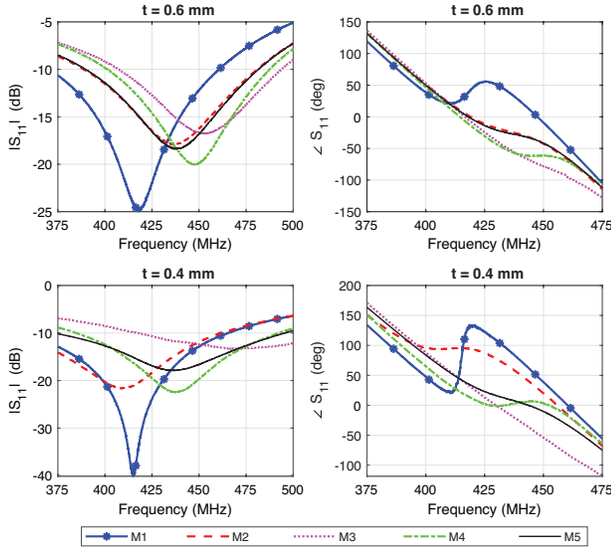


Fig. 5. Measurement results for the magnitude and the phase of the reflection coefficient obtained with the porcine model for two shell thicknesses.

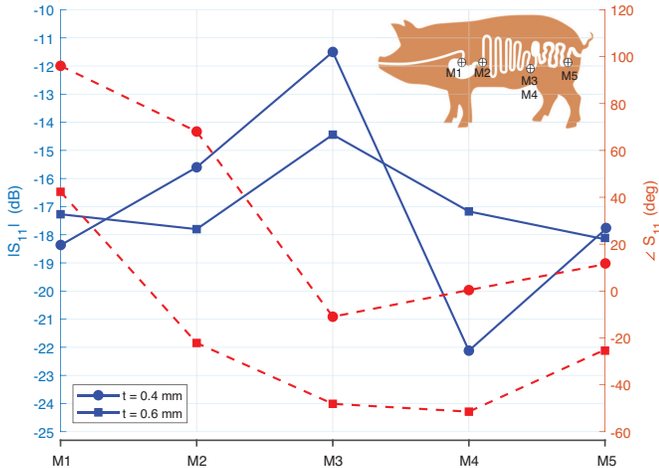


Fig. 6. The values of the magnitude and the phase of the reflection coefficient at 434 MHz measured using the porcine model. The straight lines show the magnitude and the dashed lines show the phase.

and the prototypes were inserted in 10 cm depth as shown in Fig. 3e.

Fig. 5 shows the measurement results obtained using the porcine model. As can be seen, the change in the magnitude and phase values in different tissues is greater compared to the simulations and measurements in the liquids. The responses in different tissues become more distinguishable with a more realistic setup, as the real tissues have a heterogeneous form and contain air gaps. The EM properties of the real tissues vary depending on the spatial coordinates, contrary to the homogeneous tissue models, which contributes to the differences observed in the responses.

Furthermore, Fig. 6 illustrates the magnitude and phase values measured at 434 MHz with the porcine model. It can be observed that the values change significantly depending on the tissue. This indicates that it is possible to distinguish the tissues by tracking the impedance response. Moreover,

the graphs obtained with two different antennas have similar tendencies, showing the consistency of the measurements. Hence, the results obtained with the porcine model support the idea presented in this paper.

IV. CONCLUSION

This paper examined the possibility of using the impedance detuning of ingestible antennas to track the segment of the GI tract in which a capsule is located. For this examination, meandered dipole antennas that are conformal to a PLA capsule are designed to operate in the 433 MHz ISM band. The examination is performed by comparing the impedance responses of the optimized antennas in different GI tissues with simulations and measurements (tissue-mimicking liquids and porcine model). The results show that the location of the capsule can be followed by tracking the changes in the impedance response, in phase in particular. Our future work aims at developing biodegradable resonators (for instance, Mg) on the outside part of the capsule as an additional indicator. Such a resonator will rapidly degrade in the stomach due to its acidic nature and hence can be used to distinguish the stomach from the other segments. The real-time localization of capsules inside the GI tract can be useful to improve diagnostics as well as to deliver drugs and therapies more efficiently.

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