

MedGA: A novel evolutionary method for image enhancement in medical imaging systems

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Abstract

Medical imaging systems often require the application of image enhancement techniques to help physicians in anomaly/abnormality detection and diagnosis, as well as to improve the quality of images that undergo automated image processing. In this work we introduce MedGA, a novel image enhancement method based on Genetic Algorithms that is able to improve the appearance and the visual quality of images characterized by a bimodal gray level intensity histogram, by strengthening their two underlying sub-distributions. MedGA can be exploited as a pre-processing step for the enhancement of images with a nearly bimodal histogram distribution, to improve the results achieved by downstream image processing techniques. As a case study, we use MedGA as a clinical expert system for contrast-enhanced Magnetic Resonance image analysis, considering Magnetic Resonance guided Focused Ultrasound Surgery for uterine fibroids. The performances of MedGA are quantitatively evaluated by means of various image enhancement metrics, and compared against the conventional state-of-the-art image enhancement techniques, namely, histogram equalization, bi-histogram equalization, encoding and decoding Gamma transformations, and sigmoid transformations. We show that MedGA considerably

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outperforms the other approaches in terms of signal and perceived image quality, while preserving the input mean brightness. MedGA may have a significant impact in real healthcare environments, representing an intelligent solution for Clinical Decision Support Systems in radiology practice for image enhancement, to visually assist physicians during their interactive decision-making tasks, as well as for the improvement of downstream automated processing pipelines in clinically useful measurements.

Keywords: Medical imaging systems, Image enhancement, Genetic Algorithms, Magnetic resonance imaging, Bimodal image histogram, Uterine fibroids

1. Introduction

Nowadays, medical imaging systems play a key role in the clinical workflow, thanks to their capability of representing anatomical and physiological features that are otherwise inaccessible to inspection, thus proposing accurate imaging biomarkers and clinically useful information (Rueckert et al., 2016; Lambin
5 et al., 2017). Medical images are considerably different from the pictures usually analyzed in Pattern Recognition and Computer Vision, as regards the appearance of the depicted objects as well as the information conveyed by the pixels. As a matter of fact, medical imaging techniques exploit several different principles
10 to measure spatial distributions of physical attributes of the human body, allowing us to better understand complex or rare diseases (Toennies, 2017). The effectiveness of such techniques can be reduced by a plethora of phenomena, such as noise and partial volume effects (Toennies, 2017), which might affect the measurement processes involved in imaging and data acquisition devices.
15 In addition, computer-aided medical image acquisition procedures generally include reconstruction methods (producing two, three, or even four dimensional imaging data), which could cause the appearance of artifacts. Image contrast and details might also be impaired by the procedures used in medical imaging, as well as by the physiological nature of the body part under investigation.

20 Medical images actually convey an amount of information—mainly related
to high image resolution and high pixel depth—that could overwhelm the human
vision capabilities in distinguishing among dozens of gray levels (Ortiz et al.,
2013). Improving the appearance—and the visual quality—of medical images is
therefore essential to provide physicians with valuable information that would
25 not be immediately observable in the original image, assisting them in anomaly
detection, diagnosis and treatment. This kind of diagnosis includes two basic
processes: image observation (visual perception), and diagnostic interpretation
(cognition) (Krupinski, 2010). Errors occurring in these diagnostic and thera-
peutic decision-making processes may have a significant impact on patient care,
30 most notably possible misdiagnoses. In this context, image enhancement tech-
niques aim at realizing a specific improvement in the quality of a given medical
image. The enhanced image is expected to better reveal certain features, com-
pared to their original appearance (de Araujo et al., 2014). In particular, these
methods could have a significant clinical impact when the dynamic range of the
35 actual pictorial content is not commensurable with the range of the displayed
data (i.e., monitor luminance response), as well as when the input image is char-
acterized either by a high level of noise or by a low contrast (Paranjape, 2009;
Gonzalez & Woods, 2002). This also applies to specialized computer screens for
diagnostic reporting.

40 Although the majority of the enhancement techniques are typically applied
to generate improved images for a human observer, others are exploited as a pre-
processing step to provide enhanced images to further algorithms for computer-
assisted analyses (Paranjape, 2009). The first category includes techniques de-
voted to remove noise, enhance contrast and sharpen the details. The second
45 category, partially overlapped with the former one, includes additional tech-
niques such as edge detection and object segmentation for automated process-
ing (Rangayyan, 2009). It was shown that a high-contrast medical image could
lead to a better interpretation of the different adjacent tissues in the imaged
body part (Chen et al., 2015). Accordingly, the resulting enhanced image—
50 in terms of signal intensities of different tissues—can facilitate the automated

segmentation, feature extraction, and classification of these tissues.

In the clinical routine, Contrast-Enhanced (CE) Magnetic Resonance Imaging (MRI) is a diagnostic technique that enables a more precise assessment of the imaged tissues after the administration of a Gadolinium-based contrast medium in patients (Sourbron & Buckley, 2013). MRI is currently the most prominent modality to obtain soft-tissue imaging (Brown et al., 2014), especially in oncology, since it provides significant improvements—in terms of image contrast and resolution—between lesion and healthy tissue (Metcalf et al., 2013). For these reasons, MRI is considered more suitable than Computed Tomography in determining the extent of cancer infiltration. Furthermore, the excellent MRI soft-tissue contrast has led to an increasing role of this modality in target volume delineation for therapy applications, such as image-guided surgery and radiotherapy treatment, and for patients follow-up (i.e., staging and assessing tumor response) (Evans, 2008). However, MRI data are affected by acquisition noise (Styner et al., 2000) and are also prone to imaging artifacts, related to magnetic susceptibility and large intensity inhomogeneities of the static magnetic field (i.e., streaking or shadowing artifacts (Bellon et al., 1986)), especially using high magnetic field strengths. These aspects make MR image enhancement a challenging task aiming at improving the results of automatic segmentation methods. The existing image enhancement approaches generally attempt to improve the contrast level of the whole image and do not address the issues related to overlapped gray level intensities; as a consequence, neither the region contour sharpness nor the image thresholding results can be improved. In the case of threshold-based image segmentation with two classes (i.e., foreground and background) (Muangkote et al., 2017), the input image is assumed to have a bimodal distribution of the histogram bins (Xue & Zhang, 2012). Thus, an appropriate image enhancement method that yields medical images with a sharper bimodal distribution is required. However, determining the best pre-processing of an image—able to preserve the structural information of the image, while enhancing the underlying bimodal distribution—is a complex task on a multi-modal fitness landscape that demands the use of global optimization approaches.

This paper presents a novel image enhancement technique based on Genetic Algorithms (GAs) (Holland, 1992), called MedGA, specifically aimed at strengthening the sub-distributions in medical images with an underlying bi-
85 modal histogram of the gray level intensities. Among the existing soft computing methods for global optimization, GAs represent the most suitable technique for this application, because of the discrete structure of the candidate solutions and the intrinsic combinatorial structure of the problem under investigation.

In this work, we apply MedGA to a clinical context involving CE MR image
90 analysis, i.e., Magnetic Resonance guided Focused Ultrasound Surgery (MRg-FUS) for uterine fibroids. The performances of MedGA are quantitatively evaluated by means of the most relevant image enhancement metrics, and compared against the conventional state-of-the-art image enhancement techniques, namely, histogram equalization, bi-histogram equalization, encoding and de-
95 coding Gamma transformations, and three instances of sigmoid transformation. Considering the possible clinical applications, MedGA is able to improve the visual perception of a Region of Interest (ROI) in MRI data with an underlying bimodal intensity distribution. In addition, MedGA can be used as an intelligent pre-processing step, in red any pipeline defined to realize an efficient
100 threshold-based image segmentation with two classes (i.e., binarization), applied to expert systems working on MRI data. Indeed, image thresholding approaches performed on CE MR image regions could considerably benefit from input data pre-processed by MedGA.

The main contributions of MedGA in the context of expert and intelligent
105 clinical systems can be briefly outlined as follows. MedGA acts as an expert system by playing a two-fold role: *(i)* image enhancement to visually assist physicians during their interactive decision-making tasks, and *(ii)* improvement of the results in downstream automated processing pipelines for clinically useful measurements. The rationale behind the development of MedGA is the need
110 of an intelligent model that is well-suited to effectively enhance medical images with roughly bimodal histograms. To the best of our knowledge, MedGA is the first work that explicitly deals with the improvement of thresholding-

based segmentation results (Xue & Zhang, 2012). Therefore, our computational framework can be employed as an intelligent solution in Clinical Decision Support Systems (CDSSs). As a matter of fact, MedGA represents an interpretable computational model (Castelvecchi, 2016) that allows for the understandability of the results (i.e., the gray level histogram is readable by the user). The compelling issues related to the interpretability of Machine Learning and Computational Intelligence methods in medicine are fundamental for the adoption and the clinical feasibility of a novel CDSS (Cabitza et al., 2017). In addition, no user interaction is required thanks to a reliable calibration step for the parameter settings of the GA. The goal of this paper consists in showing that evolutionary computation methods can boost the state-of-the-art performance in medical image enhancement, thus fostering GAs as a new concrete support tool for the clinical practice. Such an expert system may have a significant impact in real healthcare environments.

This manuscript is organized as follows. Related literature works and a theoretical comparison with existing methods are outlined in Section 2. Section 3 describes the MedGA image enhancement method. The evaluation metrics and the MR images used to assess the performance of MedGA are described in Section 4. Section 5 presents the achieved experimental results, by extensively explaining parameter analysis of MedGA. Finally, discussions and conclusive remarks are reported in Sections 6 and 7, respectively.

2. Background

2.1. Related work

Most of the existing enhancement techniques are empirical or heuristic methods—strongly related to a particular type of images—that generally aim at improving the contrast level of images degraded during the acquisition process (Chen et al., 2018). As a matter of fact, finding the best gray level mapping that adaptively enhances each different input image can be considered an optimization problem (Paulinas & Ušinskas, 2007; Draa & Bouaziz, 2014).

Unfortunately, no unifying theory employing a standardized image quality measure is currently available to define a general criterion for image enhancement (Munteanu & Rosa, 2004). In addition, in the case of medical imaging, techniques tailored on specific tasks are necessary to achieve a significant enhance-
145 ment and, in general, interactive procedures involving considerable human effort are needed to obtain satisfactory results.

In order to achieve objective and reproducible measurements conveying clinically useful information, operator-dependence should be minimized by means
150 of automated methods. Point-wise operations in the spatial (pixel) domain, representing the simplest form of image processing, are effective solutions since efficiency requirements have also to be met. In the case of image enhancement, they re-map each input gray level into a certain output gray level, according to a global transformation (Gonzalez & Woods, 2002). Thus, such kind of tech-
155 niques treat images as a whole, without considering specific features of different regions, or selectively distinguishing between a collection of contrast enhancement degrees or settings (Munteanu & Rosa, 2004). Histogram Equalization (HE) is the most common global image enhancement technique, whose aim is to uniformly redistribute the input gray level values according to the cumula-
160 tive density function of its histogram (Gonzalez & Woods, 2002; Hall, 1974). Unfortunately, HE does not take into account the image mean intensity (Chen & Ramli, 2003), which is subject to a significant change during the equalization process by invariably shifting the output mean brightness to the middle gray level, regardless of the mean gray level in the input image (Gan et al., 2014).
165 Consequently, HE is not able to preserve the input mean brightness, possibly suffering from over-enhancement, and giving rise to artifacts such as the so-called washed-out effect (Chen & Ramli, 2003). This global transformation method applies contrast stretching just on gray levels with the highest frequencies, causing a significant contrast loss concerning the gray levels characterized
170 by low frequencies in the input histogram (Kim, 1997). Bi-Histogram Equalization (Bi-HE), which is a refined version of the traditional HE, was proposed to overcome the limitations related to input mean brightness preservation, mainly

caused by histogram flattening (Kim, 1997). Firstly, Bi-HE splits the original histogram into two sub-histograms according to the global mean of the original image; afterwards, the sub-histograms are independently processed by applying the standard HE method to each of them.

The complexity of the enhancement criteria to be met (i.e., the effective contrast stretching combined with image detail preserving) leads to the application of global search meta-heuristics that allow for coping with several constraints, which are not generally tractable by means of traditional exhaustive computational approaches (Paulinas & Ušinskas, 2007; Munteanu & Rosa, 2004; Ortiz et al., 2013). Evolutionary methods have been widely adopted in the image enhancement domain to find the optimal enhancement kernel (Munteanu & Rosa, 2004), sequence of filters (Kohmura & Wakahara, 2006), or input-output mapping transformation (Saitoh, 1999; Carbonaro & Zingaretti, 1999). Recently, (Hashemi et al., 2010) proposed a GA-based method that efficiently encodes the histogram by means of the non-zero intensity levels, by employing genetic operators that directly process images to increase the visible details and contrast of low illumination regions, especially in the case of high dynamic ranges. The authors argued that this method yields “natural-looking” images, considering the visual appearance.

Regarding other evolutionary computation approaches, Genetic Programming (GP) (Koza, 1992) was shown to be a powerful framework to select and combine existing algorithms in the most suitable way. Differently to GAs, GP evolves a population of functions, or more generally, computer programs to solve a computational task. The solutions in the computer program space can be represented as trees, lines of code, expressions in prefix or postfix notations as well as strings of variable length (Castelli et al., 2014). For instance, (Bianco et al., 2017) tackled the video change detection problem (among the frames of video streams) by combining existing algorithms *via* different GP solutions exploiting several fusion schemes. The fitness function was composed of different performance measures regarding change detection evaluation. For what concerns the application of GP in image enhancement, (Poli & Cagnoni, 1997)

proposed an approach to yield optimally pseudo-colored images for visualization
205 purposes, aiming at combining multiple gray-scale images (e.g., time-varying
images, multi-modal medical images, and multi-band satellite images) into a
single pseudo-color image. This approach relies on user interactions to deter-
mine which candidate solution should be the winner in tournament selection,
so it does not explicitly require a fitness function. As case studies, a pair of
210 brain MRI sequences were fused as well as the motion of the heart on echocar-
diographic images was synthesized into a single pseudo-color image.

Other works exploited Swarm Intelligence techniques. The approach pre-
sented in (Shanmugavadivu & Balasubramanian, 2014), called Multi-Objective
Histogram Equalization, uses Particle Swarm Optimization (Kennedy & Eber-
215 hart, 1995) to enhance the contrast and preserve the brightness at the same
time. (Draa & Bouaziz, 2014) employed the same encoding of candidate so-
lutions and histogram mapping strategy described in (Hashemi et al., 2010),
within an optimization strategy based on the Artificial Bee Colony (ABC) al-
gorithm (Karaboga & Basturk, 2007). However, since ABC natively works in a
220 continuous space, while a discrete representation is used for the solutions (i.e.,
gray-level mapping), a discretization step is mandatory in the correction opera-
tion during the search phase. An alternative approach using the ABC algorithm
for image contrast enhancement was proposed in (Chen et al., 2018), wherein
the optimal values for the parameters of a parametric image transformation,
225 namely the Incomplete Beta Function, are estimated. Differently to the work
described in (Draa & Bouaziz, 2014), the optimization procedure is carried out
in a continuous search space. Finally, multi objective Bat Optimization and a
neuron-based model of Dynamic Stochastic Resonance were combined in (Singh
et al., 2017) for the enhancement of brain MR images.

230 *2.2. Theoretical comparison with existing methods*

It is worth noting that all works mentioned in Section 2.1 are focused on
consumer electronics or medical applications, to obtain more “visually pleasant”
images by mainly increasing the contrast of the whole image. On the contrary,

the main key novelty of MedGA consists in better revealing the two underlying
235 sub-distributions occurring in an image sub-region characterized by a roughly
bimodal histogram, overcoming the limitations of the state-of-the-art contrast
enhancement methods, which could produce false edges and consequently over-
segmentation when the input images are affected by noise, as in the case of MRI
data (Gandhamal et al., 2017). There exist other algorithms, like Histogram
240 Specification (HS), whose aim is similar to MedGA and consists in matching the
histogram of the gray level intensities of the input MR image with a desired out-
put histogram (Gonzalez & Woods, 2002). Unfortunately, this approach cannot
be applied to process image datasets characterized by a high variability in gray
level distributions, since the histogram to be matched should be defined either *a*
245 *priori* for the whole dataset, or interactively for each processed image, through
a procedure that consists in strengthening and shaping the two underlying sub-
distributions. In such cases, to automatically identify the best solution it is
more advantageous to employ global search meta-heuristics, like GAs used in
MedGA.

250 Even though MedGA exploits the same encoding of candidate solutions de-
fined in (Hashemi et al., 2010) and (Draa & Bouaziz, 2014), its purpose is very
different since it was designed to explicitly strengthen the two sub-distributions
of medical images characterized by an underlying bimodal histogram. To this
aim, we defined a specific fitness function that emphasizes the two Gaussian
255 distributions composing a bimodal histogram. This achievement plays a fun-
damental role for threshold-based segmentation approaches, since they strongly
rely on the assumption that the bimodal histogram under investigation is com-
posed of two nearly Gaussian distributions with almost equal size and variance
(Xue & Zhang, 2012).

260 MedGA also differs from GP-based approaches whose generated solutions
might have a large size (Castelli et al., 2014), even when the GP model is imple-
mented efficiently, thus representing a limitation that could significantly impair
the readability and interpretability of the final outcome. Moreover, MedGA
does not require any user interaction step, differently to (Poli & Cagnoni, 1997)

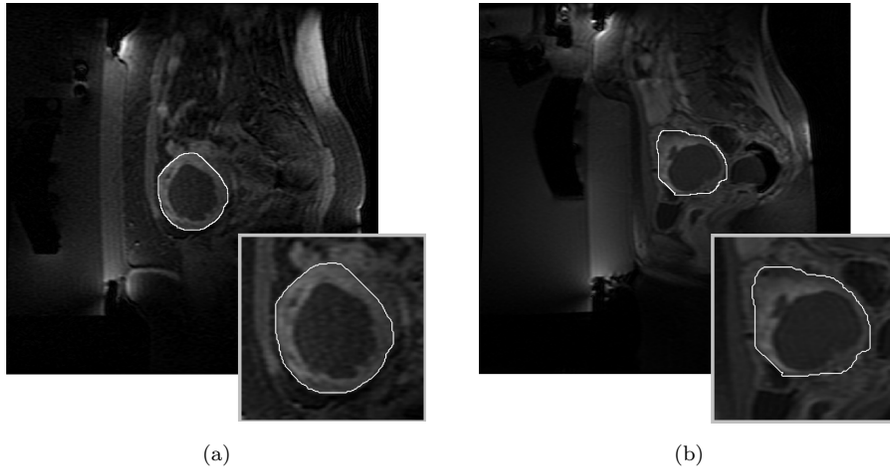


Figure 1: Examples of MR images. The ROI bounding region (i.e., the delineated uterus region), which includes the actual ROI (i.e., the uterine fibroid region), is highlighted with a white contour. The image regions including the ROIs, zoomed at the bottom right of each sub-figure, are characterized by nearly bimodal histograms.

265 where the user, being directly involved in the tournament selection, controls the evolution of simple programs that enhance and integrate multiple gray-scale images into a single pseudo-color image.

3. MedGA: an intelligent method based on Genetic Algorithms for medical image enhancement

270 MedGA is a global enhancement technique able to improve the details of medical images characterized by an underlying bimodal histogram of gray levels. Given a medical image wherein a ROI needs to be enhanced to achieve further analyses, MedGA aims at improving the ROI quality to facilitate the classification among different neighboring tissues, in order to support both the
 275 interpretation tasks by experienced radiologists and automated image analysis approaches.

The image enhancement carried out by MedGA focuses on the pixels within a sub-region of the input MRI, called ROI bounding region, including the ROI

itself. To be more precise, starting from an MR image (Fig. 1), a bounding
 280 region that roughly includes the actual ROI (e.g., the uterus region including
 uterine fibroids in Fig. 1) is identified by using either a manual or a compu-
 tational method. Afterwards, the entire original MR image is cropped at the
 smallest rectangular box including the previously identified ROI bounding re-
 gion. The pixels included in the rectangular cropped image, but external to
 285 the ROI bounding region, are set to zero (i.e., the black level). So doing, an
 image characterized by a nearly bimodal histogram is obtained. Then, a linear
 contrast stretching is applied to the initial full range of gray levels, that is, the
 ordered set $\mathcal{L}_{in} = [l_{in}^{(min)}, l_{in}^{(min)} + 1, \dots, l_{in}^{(max)} - 1, l_{in}^{(max)}] \subset \mathbb{N}$, where $l \neq l'$ for
 any $l, l' \in \mathcal{L}_{in}$. The integers $l_{in}^{(min)}$ and $l_{in}^{(max)}$ in \mathcal{L}_{in} denote the minimum and
 290 maximum non-zero gray levels of the analyzed image sub-region, respectively.
 The linear contrast stretching applied to \mathcal{L}_{in} exploits the extended range of the
 non-zero gray levels, that is, the ordered set $\mathcal{L}'_{in} = [1, \dots, l_{in}^{(max)}] \subset \mathbb{N}$, where,
 typically, $l_{in}^{(min)} > 1$. Note that the zero-padding pixels (i.e., the pixels corre-
 sponding to the level 0) are not taken into account, and that any element of \mathcal{L}_{in}
 295 is also an element of \mathcal{L}'_{in} . This normalization operation, which employs only
 values of gray levels already representable in the initial dynamic range, does not
 alter the image content and allows MedGA to process additional intensity levels
 with respect to the initial full range \mathcal{L}_{in} , by considering the intensity variability
 within the analyzed MRI dataset. It is worth noting that the pre-processing de-
 300 scribed hereby does not exploit any method that could affect the actual pictorial
 content (e.g., spatial or frequency filtering).

MedGA exploits a population P of individuals $C_i = [C_i(1), C_i(2), \dots, C_i(n)]$
 (with $i = 1, \dots, |P|$) defined as circular arrays of integer numbers of size n ,
 where $n = |\mathcal{L}'_{in}|$ corresponds to the number of different gray levels belonging to
 305 \mathcal{L}'_{in} identified in the input MR image (i.e., the gray levels whose frequency is
 greater than zero in the input MR image). Each individual $C_i \in P$ is randomly
 initialized by sampling n integer values from the discrete uniform distribution
 in \mathcal{L}'_{in} . The n values are then sorted in ascending order so that the intensity
 levels $C_i(j)$ (with $j = 1, \dots, n$) codified by the individual can be mapped to

310 the intensity levels of the input MR image (i.e., the gray level frequencies of
the input MR image are assigned to the corresponding intensity levels of the
individual). During the initialization of the individuals, if an integer value is
sampled more than once then the frequency values of the input MR image,
corresponding to these gray level intensities, are summed up and assigned to
315 the same gray level of the individual.

The rationale of MedGA is to process the ordered set \mathcal{L}'_{in} by modifying its
gray levels using a sequence of genetic operators, to obtain a solution character-
ized by a stronger bimodal gray level distribution in an output gray-scale range
 $\mathcal{L}_{out} = [l_{out}^{(min)}, \dots, l_{out}^{(max)}] \subset \mathbb{N}$, where any element of \mathcal{L}_{out} is also an element of
320 \mathcal{L}'_{in} . In such a way, a direct mapping between the gray levels of the original im-
age and the final one is defined, so that each gray level in the original histogram
is replaced with the gray level value contained by the same position in the final
best solution $C_{best} \in P$. MedGA realizes this global intensity transformation
by improving the separation between $\mathcal{H}_{1,i}$ and $\mathcal{H}_{2,i}$, which represent the dark
325 and bright sub-regions of the histogram \mathcal{H}_i encoded by the individual C_i , re-
spectively. To this aim, MedGA exploits the optimal threshold $\theta_{opt,i}$ adaptively
selected using the Iterative Optimal Threshold Selection (IOTS) method (Ridler
& Calvard, 1978; Trussell, 1979). This procedure yields enhanced medical im-
ages that better reveal the bimodal intensity distribution in computer-assisted
330 ROI extraction tasks.

At each iteration of MedGA, a number of individuals properly selected from
the current population are inserted into intermediate populations, and modified
by means of crossover and mutation operators. Note that, at each iteration,
each individual C_i belonging to the current population codifies for an ordered
335 set $\mathcal{L}_{out,i} = [C_i(1), \dots, C_i(n)] = [l_{out,i}^{(min)}, \dots, l_{out,i}^{(max)}]$, which represents a modi-
fied gray level distribution of \mathcal{L}'_{in} . In order to simplify the notation, in what
follows we do not explicitly express that, *at each iteration*, each individual C_i
is (possibly) represented by a different circular array corresponding to \mathcal{L}_{out} .

For the selection of individuals, MedGA exploits a *tournament* strategy for
340 three main reasons: (*i*) the selection pressure can be controlled by setting the

tournament size k (with $k \ll |P|$); (ii) the fitness evaluations are performed only on the k individuals selected for the tournaments, and not on the whole population; (iii) this technique could be easily implemented on parallel architectures (Miller & Goldberg, 1995).

345 A *single point crossover* operator is applied with a given probability p_c to the individuals selected by the tournament strategy and belonging to the first intermediate population P' . Namely, given two parent individuals $C_m, C_f \in P'$ (for some $m, f = 1, \dots, |P|$), a crossover point is randomly selected from the circular arrays $[C_m(1), C_m(2), \dots, C_m(n)]$ and $[C_f(1), C_f(2), \dots, C_f(n)]$ encoding the individuals, and two offspring are generated by swapping 50% of the values between the two parents. The offspring are then inserted into a second intermediate population P'' .

The *mutation* operator is applied with probability p_m to each element $C_i(j) \in C_i = [l_{out,i}^{(min)}, \dots, l_{out,i}^{(max)}]$ of each individual belonging to P'' , where $l_{out,i}^{(min)}$ and 355 $l_{out,i}^{(max)}$ are the minimum and maximum non-zero gray levels encoded by C_i during the current iteration, respectively. In particular, if the gray level intensity encoded in $C_i(j)$ is smaller than the optimal threshold $\theta_{opt,i}$ evaluated by IOTS at that iteration for the individual C_i , then an integer is randomly sampled from the uniform distribution in $[l_{out,i}^{(min)}, \dots, \theta_{opt,i} - 1] \subset \mathbb{N}$ to update the value 360 $C_i(j)$; otherwise, an integer is randomly sampled from the uniform distribution in $[\theta_{opt,i}, \dots, l_{out,i}^{(max)}] \subset \mathbb{N}$ to update the value $C_i(j)$.

Finally, to prevent the quality of the best solution from decreasing during the optimization, MedGA also exploits an *elitism* strategy, so that the best individual from the current population is copied into the next population without 365 undergoing the genetic operators.

Fig. 2 illustrates the overall procedure of MedGA, by presenting the initialization phase as well as the flow diagram of the proposed GA for image enhancement. The final best solution shows the achieved result on MRI data characterized by a bimodal histogram, emphasizing the two underlying distributions for the subsequent image thresholding phase, according to the optimal 370 adaptive threshold θ_{opt} , computed on the final best solution C_{best} , by using the

simple IOTS algorithm (Ridler & Calvard, 1978; Trussell, 1979).

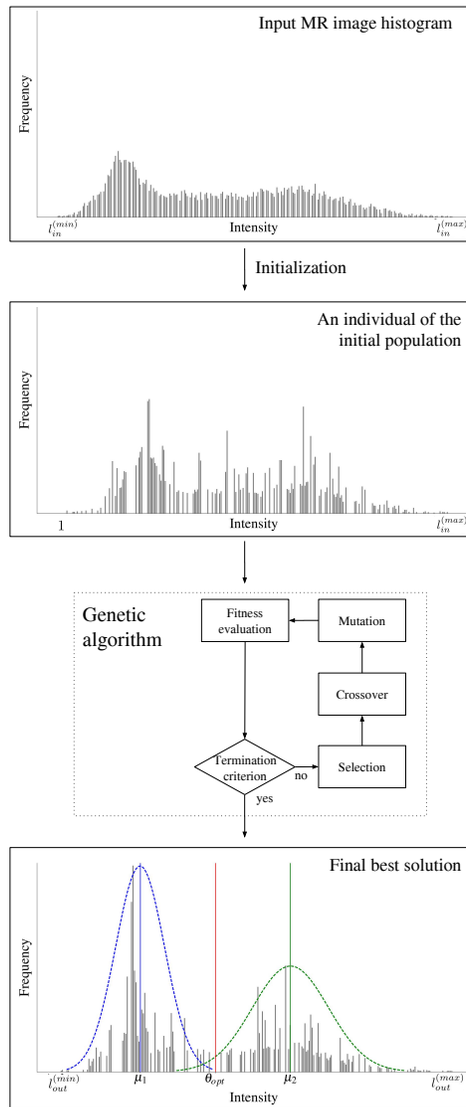


Figure 2: Workflow of MedGA: the individuals are initialized according to the characteristics of the input MR image, and processed by the GA. The final best solution of MedGA strengthens the two underlying distributions in the gray levels intensity characterized by mean values μ_1 and μ_2 and standard deviations σ_1 and σ_2 , respectively. The two distributions are highlighted in the plot with blue and green dashed lines.

To evaluate the quality of the individuals throughout the optimization, the fitness function has been conceived to obtain a bimodal histogram in the gray levels intensities, therefore facilitating further automated image processing phases. The fitness function $\mathcal{F}(\cdot)$ used by MedGA fosters individuals C_i characterized by a histogram with two well separated normal distributions, having an equal distance from the optimal threshold $\theta_{opt,i}$. To this purpose, at each iteration MedGA estimates, by using the efficient IOTS algorithm (Ridler & Calvard, 1978; Trussell, 1979), the mean values $\mu_{1,i}$ and $\mu_{2,i}$ of the two components $\mathcal{H}_{1,i}$ and $\mathcal{H}_{2,i}$ of the histogram \mathcal{H}_i , encoded by each individual C_i , according to the current optimal threshold $\theta_{opt,i}$. Afterwards, at each iteration the fitness function of every individual C_i is calculated as follows:

$$\begin{aligned}
\mathcal{F}(C_i) &= \tau_1 + \tau_2 + \tau_3, \quad \text{where:} \\
\tau_1 &= |2 \cdot \theta_{opt,i} - \mu_{1,i} - \mu_{2,i}|, \\
\tau_2 &= |\omega_{1,i} - 3\sigma_{1,i}|, \\
\tau_3 &= |\omega_{2,i} - 3\sigma_{2,i}|.
\end{aligned} \tag{1}$$

The terms $\omega_{1,i} = \frac{1}{2}(\theta_{opt,i} - \min_{j \in \{1, \dots, n\}} \{C_i(j)\})$ and $\omega_{2,i} = \frac{1}{2}(\max_{j \in \{1, \dots, n\}} \{C_i(j)\} - \theta_{opt,i})$ correspond to the half width of $\mathcal{H}_{1,i}$ and $\mathcal{H}_{2,i}$, respectively, while $\sigma_{1,i}$ and $\sigma_{2,i}$ are the standard deviations of $\mathcal{H}_{1,i}$ and $\mathcal{H}_{2,i}$, respectively. The three terms of the fitness function $\mathcal{F}(\cdot)$ cooperate together to achieve the desired image enhancement: τ_1 aims at maintaining the mean values $\mu_{1,i}$ and $\mu_{2,i}$ equidistant from the yielded optimal threshold $\theta_{opt,i}$, while τ_2 and τ_3 are designed to force the sub-histograms $\mathcal{H}_{1,i}$ and $\mathcal{H}_{2,i}$ to approximate normal distributions.

We exploit the empirical property of normal distributions related to the coverage probability with respect the standard deviation. To be more precise, we consider the so-called 3- σ rule, which states that approximately 99.73% of the values lie within 3σ according to: $Pr(\mu - 3\sigma \leq X \leq \mu + 3\sigma) \approx 0.9973$, where μ , σ and X represent the mean, the standard deviation and an observation from a normally distributed random variable, respectively.

Examples of MR image enhancement results, achieved by MedGA on uterine fibroid, are shown in Fig. 3. MedGA enhances the input MR image by making

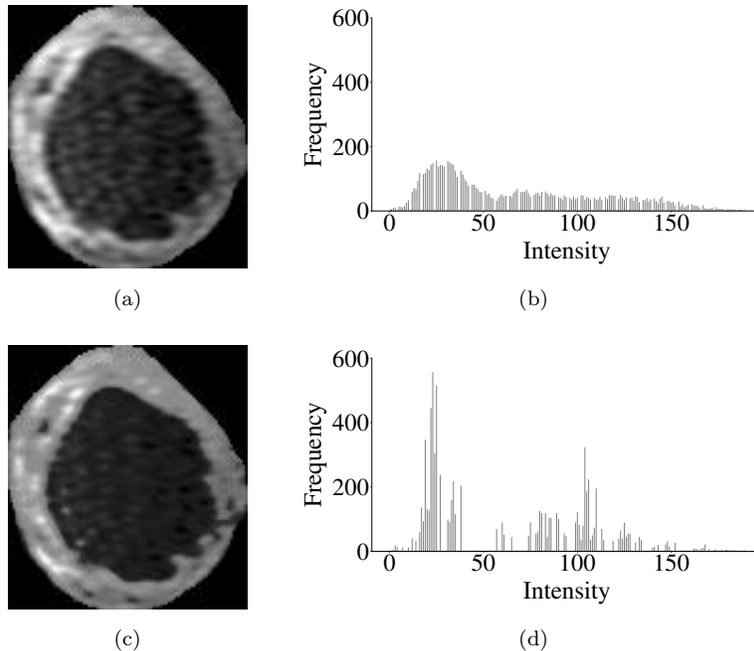


Figure 3: Enhanced image obtained by MedGA on uterine fibroids of patients who had undergone MRgFUS therapy: (a) input MR image normalized by using linear contrast stretching on the initial full range of the masked MR image; (c) output image after the application of MedGA. Plots (b) and (d) report the histograms of gray level intensities corresponding to the MR images in (a) and (c), respectively.

uterine fibroid regions more uniform and with strong edges in terms of both visual human perception and automated image processing. The histogram in
 390 Fig. 3d proves that the output image is characterized by a more defined bimodal distribution compared to the initial image (Fig. 3b), which roughly tends to a trimodal gray level distribution.

It is worth noting that, in the case of further automated analysis, ROI pixel classification can be carried out by means of a basic threshold-based segmenta-
 395 tion approach, since MR images enhanced with MedGA reveal a more precise separation between the two (possibly overlapping) sub-distributions in the histogram. Accordingly, MedGA allows for dealing with image histograms that do not meet the assumptions imposed by thresholding techniques, regarding bi-

modal histograms composed of two nearly Gaussian distributions with almost
400 equal size and variance (Xue & Zhang, 2012). Indeed, MedGA enhances the
image thresholding results on MRI data, as shown in Fig. 2, where the final
best solution improves the underlying bimodal nature of the input histogram.

A sequential and a parallel version of MedGA have been implemented. The
sequential version has been entirely developed using the Python programming
405 language (version 2.7.12), while the parallel version is based on a Master-Slave
paradigm employing `mpi4py`, which provides bindings of the Message Passing
Interface (MPI) specifications for Python to leverage High-Performance Com-
puting (HPC) resources (Dalcín et al., 2005).

4. Materials and evaluation metrics

4.1. MRI dataset

Eighteen patients affected by symptomatic uterine fibroids, who underwent
MRgFUS therapy, were taken into account. The total number of the exam-
ined fibroids was 29 represented on 163 CE MR slices, since some patients
presented a pathological scenario with multiple fibroids. The analyzed images
415 were acquired using a Signa HDxt MRI scanner (General Electric Medical Sys-
tems, Milwaukee, WI, USA) at two different institutions. These MRI data
were acquired after the MRgFUS treatment, executed with the ExAblate 2100
(Insightec Ltd., Carmel, Israel) HIFU equipment. The considered MRI series
were scanned using the T1w “Fast Spoiled Gradient Echo + Fat Suppression +
420 Contrast mean” (FSPGR+FS+C) protocol. This MRI protocol is usually em-
ployed for Non-Perfused Volume (NPV) assessment (Gorny et al., 2011), since
ablated fibroids appear as hypo-intense areas due to low perfusion of the con-
trast medium. Sagittal MRI sections were processed, in compliance with the
current clinical practice for therapy response assessment (Militello et al., 2014).
425 MRI acquisition parameters were: Repetition Time (TR): 150-260 ms; Echo
Time (TE): 1.392-1.544 ms; pixel depth: 16-bit; matrix size: 512×512 pixels;

slice thickness: 5.0 mm; slice spacing: 6.0 mm; pixel spacing: 0.6641-0.7031 mm.

As explained in Section 3, before applying MedGA image enhancement, the
 430 uterus region (i.e., ROI bounding region) must be delineated. This task can be
 accomplished manually by the user or automatically by means of computational
 methods to reduce operator-dependency, as previously proposed in (Militello
 et al., 2015b). Afterwards, the pixels with values lower than the optimal thresh-
 old θ_{opt} , computed by means of the efficient IOTS method (Ridler & Calvard,
 435 1978; Trussell, 1979), are segmented in the binarized MR image. In this clinical
 scenario, segmentation approaches must deal with NPV inhomogeneities, due
 to sonication spots during the MRgFUS treatment.

4.2. Image enhancement evaluation metrics

In this section, we recall the definition of the metrics typically used to eval-
 440 uate image enhancement approaches, which will be exploited to assess the per-
 formance of MedGA. These metrics are essential to quantitatively evaluate the
 effects of image enhancement techniques, since measuring the “quality” of an
 image might be strongly subjective. In particular, we benefit here from the
 metrics considered in (Hashemi et al., 2010) to assess the capability of image
 445 enhancement approaches in improving contrast, details and human visual per-
 ception.

Let I_{orig} and I_{enh} be the original input image and the enhanced image,
 respectively, consisting in M rows and N columns. Considering that the range of
 gray levels of the output image $\mathcal{L}_{out} = [l_{out}^{(min)}, \dots, l_{out}^{(max)}]$ could be different from
 the original range $\mathcal{L}_{in} = [l_{in}^{(min)}, \dots, l_{in}^{(max)}]$, as a first step before computing the
 metrics we remap the output pixel intensities (i.e., the gray levels) into the
 original range, as follows:

$$\tilde{I}_{enh}(a, b) = \frac{(I_{enh}(a, b) - l_{out}^{(min)}) \cdot (l_{in}^{(max)} - l_{in}^{(min)})}{(l_{out}^{(max)} - l_{out}^{(min)})} + l_{in}^{(min)}, \quad (2)$$

where $a = 1, \dots, M$ and $b = 1, \dots, N$. Note that we actually consider the
 extended range \mathcal{L}'_{in} for I_{orig} , as explained in Section 3.

The Peak Signal-to-Noise Ratio (*PSNR*) denotes the ratio between the maximum possible intensity value of a signal and the distortion between the input and output images:

$$\begin{aligned} PSNR &= 10 \cdot \log_{10} \left(\frac{(l_{in}^{(max)})^2}{MSE} \right) \\ &= 20 \cdot \log_{10} \left(\frac{l_{in}^{(max)}}{\sqrt{MSE}} \right), \end{aligned} \quad (3)$$

where $MSE = \frac{1}{M \times N} \sum_{a=1}^M \sum_{b=1}^N \|I_{orig}(a, b) - \tilde{I}_{enh}(a, b)\|^2$ is the Mean Squared Error, which allows us to compare the pixel values of I_{orig} to those of \tilde{I}_{enh} .

Furthermore, the *PSNR* is usually expressed in terms of the logarithmic Decibel scale. With regard to our application, we employ only a limited portion of the full dynamic range of the 16-bit images (see Section 3); we thus use as the largest possible value the maximum intensity value present in the original image (i.e., $l_{in}^{(max)} = \max\{\mathcal{L}_{in}\} = \max\{\mathcal{L}'_{in}\}$) instead of the maximum representable value in a 16-bit image (i.e., $2^{16} - 1 = 65,535$).

Munteanu & Rosa (2004) stated that good contrast and enhanced images are characterized by high numbers of *edgels* (i.e., pixels belonging to an edge), and that an enhanced image should have a higher intensity of the edges, compared to its non-enhanced counterpart (Saitoh, 1999). Therefore, a good enhancement technique should yield satisfactory results in the case of standard vision processing tasks, such as segmentation or edge detection (Starck et al., 2003). Here, to evaluate image enhancement of MRI data, we employ the method proposed by Canny (1986), which is a highly reliable and mathematically well-defined edge detector. This approach deals with weak edges and accurately determines edgels, by applying a double threshold (to identify potential edges) and a hysteresis-based edge tracking. Let \mathcal{M}_{Canny} be the edge map yielded by the Canny’s edge detector, which is a binary image wherein only edgels are set to 1. The number of detected edges (*#DE*) in \mathcal{M}_{Canny} is computed as:

$$\#DE = \sum_{a=1}^M \sum_{b=1}^N \mathcal{M}_{Canny}(a, b). \quad (4)$$

An additional metrics, called Absolute Mean Brightness Error (*AMBE*) (Chen & Ramli, 2003; Arriaga-Garcia et al., 2014), can be employed to measure the brightness preservation of the enhanced image:

$$AMBE = \frac{|\mathbb{E}[I_{orig}] - \mathbb{E}[\tilde{I}_{enh}]|}{L}, \quad (5)$$

where $\mathbb{E}[\cdot]$ denotes the expected (mean) value of a gray level distribution. *AMBE* is normalized in $[0, 1]$, divided by $L = l_{in}^{(max)} - l_{in}^{(min)}$, which is the dynamic range of the input gray-scale (in our case, \mathcal{L}'_{in}). Note that low values of *AMBE* denote
460 that the mean brightness of the original image is preserved.

Finally, we consider an alternative quality metrics called Structural Similarity Index (*SSIM*) (Wang et al., 2004), used to assess the image degradation perceived as variations in structural information (Bhandari et al., 2016). The structural information defines the attributes that represent the structure of ob-
465 jects in the image, independently of the average luminance and contrast. In particular, local luminance and contrast are taken into account since overall values of luminance and contrast can remarkably vary across the whole image. *SSIM* is based on the degradation of structural information—assuming that human visual perception is highly adapted for extracting structural information
470 from a scene—and compares local patterns of pixel intensities. As a matter of fact, natural image signals are highly structured, since pixels are strongly dependent on each other, especially those close by. These dependencies convey important information about the structure of the objects in the viewing field. Let \mathbf{X} and \mathbf{Y} be the I_{orig} and I_{enh} image signals, respectively; *SSIM* combines
475 three relatively independent terms:

- the luminance comparison $l(\mathbf{X}, \mathbf{Y}) = \frac{2\mu_{\mathbf{X}}\mu_{\mathbf{Y}} + \kappa_1}{\mu_{\mathbf{X}}^2 + \mu_{\mathbf{Y}}^2 + \kappa_1}$;
- the contrast comparison $c(\mathbf{X}, \mathbf{Y}) = \frac{2\sigma_{\mathbf{X}}\sigma_{\mathbf{Y}} + \kappa_2}{\sigma_{\mathbf{X}}^2 + \sigma_{\mathbf{Y}}^2 + \kappa_2}$;
- the structural comparison $s(\mathbf{X}, \mathbf{Y}) = \frac{\sigma_{\mathbf{X}\mathbf{Y}} + \kappa_3}{\sigma_{\mathbf{X}}\sigma_{\mathbf{Y}} + \kappa_3}$;

where $\mu_{\mathbf{X}}$, $\mu_{\mathbf{Y}}$, $\sigma_{\mathbf{X}}$, $\sigma_{\mathbf{Y}}$, and $\sigma_{\mathbf{X}\mathbf{Y}}$ are the local means, standard deviations, and cross-covariance for the images \mathbf{X} and \mathbf{Y} , while $\kappa_1, \kappa_2, \kappa_3 \in \mathbb{R}^+$ are regu-

larization constants for luminance, contrast, and structural terms, respectively, exploited to avoid instability in the case of image regions characterized by local mean or standard deviation close to zero. Typically, small non-zero values are employed for these constants; according to Wang et al. (2004), an appropriate setting is $\kappa_1 = (0.01 \cdot L)^2$, $\kappa_2 = (0.03 \cdot L)^2$, $\kappa_3 = \kappa_2/2$, where L is the dynamic range of the pixel values in I_{orig} (represented in \mathcal{L}'_{in}). $SSIM$ is then computed by combining the components described above:

$$SSIM = l(\mathbf{X}, \mathbf{Y})^\alpha \cdot c(\mathbf{X}, \mathbf{Y})^\beta \cdot s(\mathbf{X}, \mathbf{Y})^\gamma, \quad (6)$$

where $\alpha, \beta, \gamma > 0$ are weighting exponents. As reported in Wang et al. (2004), if $\alpha = \beta = \gamma = 1$ and $\kappa_3 = \kappa_2/2$, the $SSIM$ becomes:

$$SSIM = \frac{(2\mu_{\mathbf{X}}\mu_{\mathbf{Y}} + \kappa_1)(2\sigma_{\mathbf{X}\mathbf{Y}} + \kappa_2)}{(\mu_{\mathbf{X}}^2 + \mu_{\mathbf{Y}}^2 + \kappa_1)(\sigma_{\mathbf{X}}^2 + \sigma_{\mathbf{Y}}^2 + \kappa_2)}. \quad (7)$$

$SSIM$ generalizes the Universal Quality Index (UQI), defined in Wang & Bovik
480 (2002), which is obtained by setting $\kappa_1 = \kappa_2 = 0$, and yields unstable results when either $(\mu_{\mathbf{X}}^2 + \mu_{\mathbf{Y}}^2)$ or $(\sigma_{\mathbf{X}}^2 + \sigma_{\mathbf{Y}}^2)$ tends to zero. Notice that the higher the $SSIM$ value, the higher the structural similarity, implying that the enhanced image I_{enh} and the original image I_{orig} are quantitatively similar.

5. Results

485 This section presents the experimental results achieved by our image enhancement method. We first analyze the performances of MedGA by varying the parameter settings of the GA at the basis of our methodology; we then compared it against the most common and popular image enhancement techniques in the image processing field (see Gonzalez & Woods (2002) for additional
490 details).

5.1. MedGA settings analysis and calibration

To analyze the performances of MedGA and identify the best settings for the image enhancement problem, we considered a calibration set consisting of

80 medical images randomly selected from the available MRI data (described in
 495 Section 4.1), and we varied the settings of MedGA used throughout the opti-
 mization process, that is: (i) the size of the population $|P| \in \{50, 100, 150, 200\}$;
 (ii) the crossover probability $p_c \in \{0.8, 0.85, 0.9, 0.95, 1.0\}$; (iii) the mutation
 probability $p_m \in \{0.01, 0.05, 0.1, 0.2\}$; (iv) the size of the tournament selection
 strategy $k \in \{5, 10, 15, 20\}$. In all tests MedGA was run for $T = 100$ iterations.
 500 Each MedGA execution was performed by varying one setting at a time, for a
 total of 320 different settings tested and a total number of $320 \times 80 = 25600$
 MedGA executions.

The results of these tests (data not shown) highlighted that, for each value
 of $|P|$, the best settings in terms of fitness values achieved are:

- 505 1. $|P| = 50, p_c = 0.85, p_m = 0.01, k = 15$;
2. $|P| = 100, p_c = 0.9, p_m = 0.01, k = 20$;
3. $|P| = 150, p_c = 0.85, p_m = 0.01, k = 20$;
4. $|P| = 200, p_c = 0.85, p_m = 0.01, k = 20$.

Figure 4 reports the comparison of the performances achieved by MedGA
 510 with these settings, where the Average Best Fitness (ABF) was computed by
 taking into account, at each iteration of MedGA, the fitness value of the best
 individuals over the 80 optimization processes. It is clear from the plot that,
 despite the final ABF values are comparable in all settings, the convergence
 speed increases with the size of the population, as well as the running time
 required by MedGA; therefore, to choose the best settings, we analyzed the
 515 computational performances concerning the 4 tests listed above.

Considering that an MRI series related to a single patient contains on av-
 erage 10 slices with ROI fibroids, we tested the clinical feasibility of MedGA
 by calculating the total execution time for enhancing 10 randomly chosen MR
 520 images. For what concerns the tests 1-4 described above, the executions lasted
 on average 672.12 s, 1290.15 s, 1987.8 s, and 2669.74 s, respectively, for the
 optimization of the same batch of 10 images running a single core of the Intel
 Xeon E5-2440 CPU with 2.40 GHz clock frequency. On the other hand, by ex-

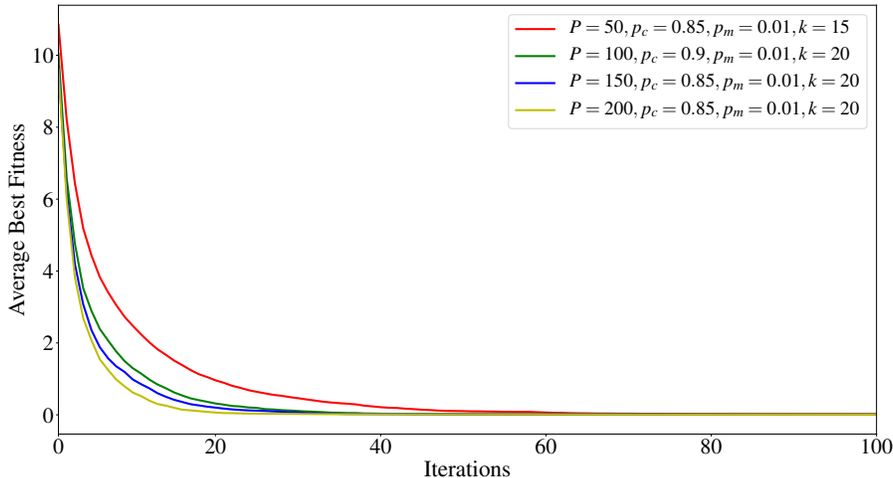


Figure 4: Comparison of the ABF achieved by MedGA with the best parameterizations found for each value of $|P|$ tested here. The average was computed over the results of the optimization of 80 MR images.

exploiting the 6 cores of the same CPU to execute the parallel version of MedGA,
 525 we achieved up to $3.6\times$ speed-up with respect to the sequential version. The
 results achieved using the parallel version of MedGA confirm the importance
 of HPC solutions in the field of real healthcare environment to obtain clinically
 feasible outcomes, that is, enhancing MR images in reasonable time for medical
 imaging practice.

530 By considering both the performance of MedGA in terms of ABF and the
 running time required to process 80 images, we selected the parameter settings
 $|P| = 100, p_c = 0.9, p_m = 0.01, k = 20$ as the best trade-off characterized by a
 good convergence speed and an adequate running time (for this specific appli-
 cation), and we exploited this configuration for all tests reported and discussed
 535 in the following section.

5.2. Comparison with state-of-the-art methods

The performances of MedGA were compared against the following image
 enhancement techniques:

- 540
 • Histogram Equalization (HE) (Paranjape, 2009; Hall, 1974), which adjusts pixel intensities for contrast enhancement according to the normalized histogram of the original image I_{orig} . With HE, gray levels are more uniformly distributed on the histogram, by spreading the most frequent intensity values;
- 545
 • Bi-Histogram Equalization (Bi-HE) (Kim, 1997)—a modification of the traditional HE—that addresses issues concerning mean brightness preservation;
- 550
 • Gamma Transformation (GT), which is a non-linear operation using the power-law relationship $s(r) = cr^\gamma$, where r and s are the input and the output gray-scale values, respectively, and c is a multiplication constant ($c = 1$ in the following tests). The parameter γ is set to values greater than 1 (i.e., decoding gamma) to obtain a gamma expansion, or to values smaller than 1 (i.e., encoding gamma) to realize a gamma compression (see Fig. 5a). In our tests we considered the values $\gamma = 0.4$ and $\gamma = 2.5$, as higher (lower) values of γ tend to logarithmic (anti-logarithmic) functions, resulting in an excessively bright (dark) output image, unsuitable for practical medical applications (Gandhamal et al., 2017);
- 555
 • Sigmoid intensity Transformation (ST) function (Fig. 5b), also called S-shaped curve, which is a global non-linear mapping defined as follows:

$$s(r) = \frac{l_{in}^{(max)}}{1 + \exp(-\lambda(r - \alpha))}, \quad (8)$$

560
 where $l_{in}^{(max)} = \max\{\mathcal{L}_{in}\} = \max\{\mathcal{L}'_{in}\}$ is the asymptotic maximum value of the function, $\alpha = \frac{1}{2} \left(l_{in}^{(max)} - l_{in}^{(min)} \right)$ is the midpoint value, and λ defines the function steepness. This transformation stretches the intensity around the level α , by making the hypo-intense histogram part darker and the hyper-intense histogram part brighter. Thus, the difference between the minimum and maximum gray values and the gradient magnitude of the image are increased, obtaining strong edges (Gandhamal et al., 2017).

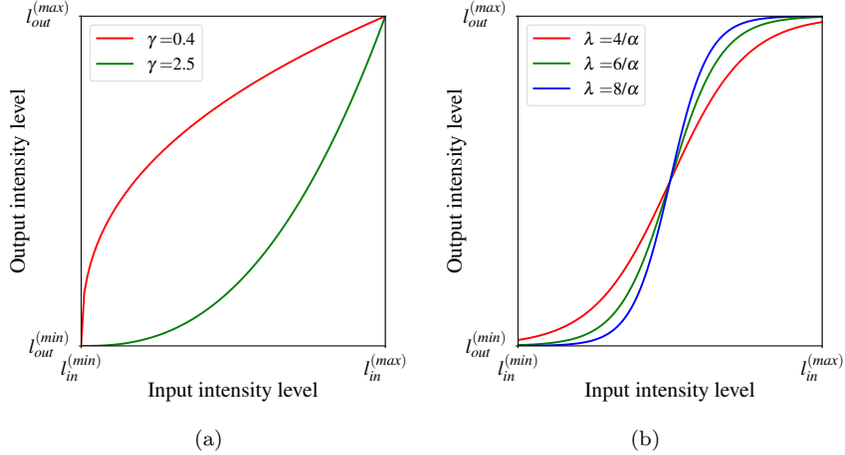


Figure 5: Plots of the implemented global non-linear intensity transformations for image enhancement: (a) Gamma Transformation; (b) Sigmoid intensity Transformation. We report on the x -axis the input intensity range $[l_{in}^{(min)}, \dots, l_{in}^{(max)}]$, and on the y -axis the output intensity range $[l_{out}^{(min)}, \dots, l_{out}^{(max)}]$.

In our tests, we used sigmoid functions that allow for considering the entire input dynamic range, by varying the curve slope with the values

565 $\lambda \in \left\{ \frac{4}{\alpha}, \frac{6}{\alpha}, \frac{8}{\alpha} \right\}$.

In order to achieve a thorough comparison between MedGA and the enhancement techniques listed above, we exploited the entire set of MRI data consisting in 163 medical images of uterine fibroids (including both calibration

570 and unseen data). For each technique, we computed the metrics $PSNR$, $\#DE$, $AMBE$, $SSIM$ defined in Section 4.2. The resulting values are given in Fig. 6 (boxplots) and in Table 1 (median, mean and standard deviation). Overall, these results highlight that MedGA significantly outperforms the conventional image enhancement approaches in terms of signal quality and perceived structural information in the images, while preserving the input mean brightness.

575 As a matter of fact, MedGA achieves the highest $PSNR$ and $SSIM$ median and mean values (see Fig. 6 and Table 1). Concerning the other enhancement techniques, we observe that Bi-HE achieves better performances with respect to

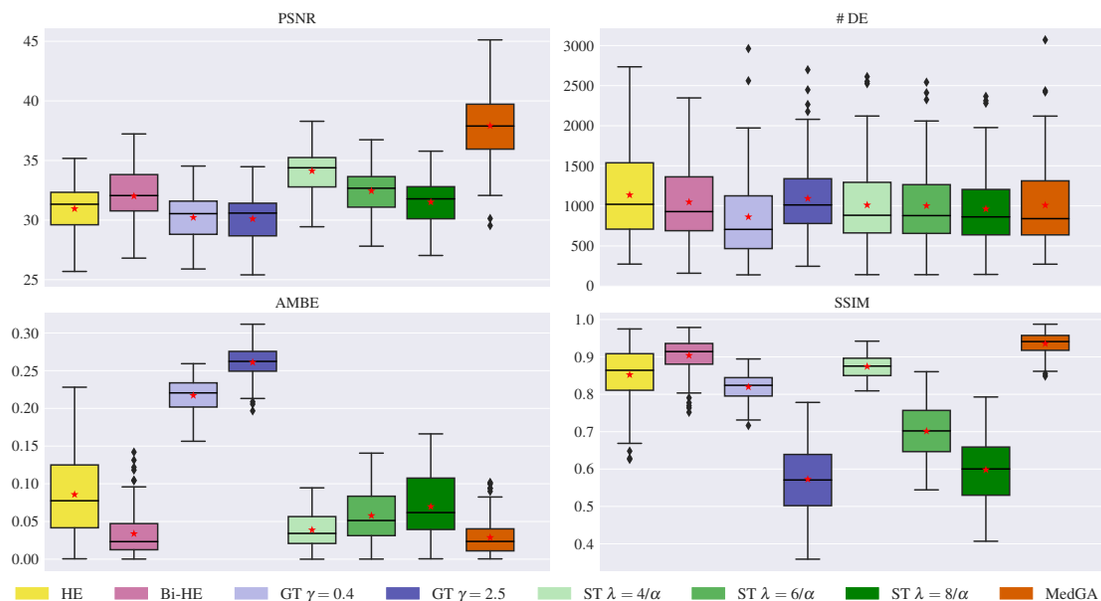


Figure 6: Boxplots of the image enhancement evaluation metrics considered in this work, obtained on an MRI dataset composed of 18 patients with uterine fibroids who had undergone MRgFUS treatment (total number of MR slices: 163). The lower and the upper bounds of each box represent the first and third quartiles of the statistical distribution, respectively, corresponding to the interquartile range. The median and the mean values are represented by a black solid line and a red star, respectively. Whiskers value is 1.5 in all cases and outliers are displayed as black diamonds.

Table 1: Median, mean and standard deviation of image enhancement evaluation metrics achieved by HE, Bi-HE, GT (with $\gamma \in \{0.4, 2.5\}$), ST (with $\lambda \in \{4/\alpha, 6/\alpha, 8/\alpha\}$) and MedGA, considering the uterine fibroid MRI dataset consisting in a total of 163 medical images.

	PSNR			#DE			AMBE			SSIM		
	Median	Mean	Std. Dev.	Median	Mean	Std. Dev.	Median	Mean	Std. Dev.	Median	Mean	Std. Dev.
HE	31.331	30.957	2.068	1020	1136.865	561.791	0.077	0.086	0.053	0.864	0.852	0.072
Bi-HE	32.059	32.014	2.232	929	1048.301	472.550	0.023	0.034	0.029	0.914	0.904	0.046
GT $\gamma = 0.4$	30.540	30.223	1.994	707	862.871	510.196	0.221	0.217	0.022	0.824	0.820	0.033
GT $\gamma = 2.5$	30.588	30.106	2.063	1012	1092.847	444.502	0.262	0.261	0.021	0.571	0.573	0.094
ST $\lambda = 4/\alpha$	34.389	34.127	1.880	883	1011.061	468.799	0.034	0.039	0.024	0.875	0.874	0.031
ST $\lambda = 6/\alpha$	32.673	32.449	1.950	879	1002.196	443.344	0.051	0.058	0.035	0.702	0.701	0.074
ST $\lambda = 8/\alpha$	31.782	31.513	1.983	863	962.141	411.109	0.062	0.070	0.042	0.600	0.598	0.089
MedGA	37.891	37.914	2.821	841	1008.773	499.906	0.024	0.029	0.023	0.941	0.936	0.029

HE, while increasing the values of γ and λ corresponds to a degradation of the
580 performances of GT and ST, respectively.

For what concerns the #DE metrics, HE over-enhances the processed MR images, as denoted by the highest median and mean values achieved, while Bi-HE allows for the preservation of the mean brightness, as also indicated by the lowest mean value of *AMBE*. On the contrary, GT achieved the worst
585 performance in terms of *AMBE* metrics. This poor result is due to the intrinsic nature of GT that transforms the input gray-scale range into a darker ($\gamma > 1$) or brighter ($\gamma < 1$) one, by increasing the number of hyper-intense and hypo-intense pixels, respectively. By doing so, GT does not preserve the input mean brightness, thus obtaining the highest values of *AMBE*. All the other methods
590 are characterized by comparable #DE values, while the *PSNR* is characterized by very high mean values in the case of GT with respect to the other techniques. On the one hand, considering the *SSIM* metrics, GT with $\gamma = 0.4$ remarkably yields better results compared to GT with $\gamma = 2.5$, especially in the case of the *SSIM*; on the other hand, all metrics related to the tested ST functions show
595 that their performances decrease as the value of λ increases. This phenomenon is related to the rapid variation characterizing the highest values of λ , which do not allow for taking into consideration the existing dependency among the pixels, especially those in the neighborhood.

Finally, we show in Fig. 7 two examples of MR images enhanced by using all

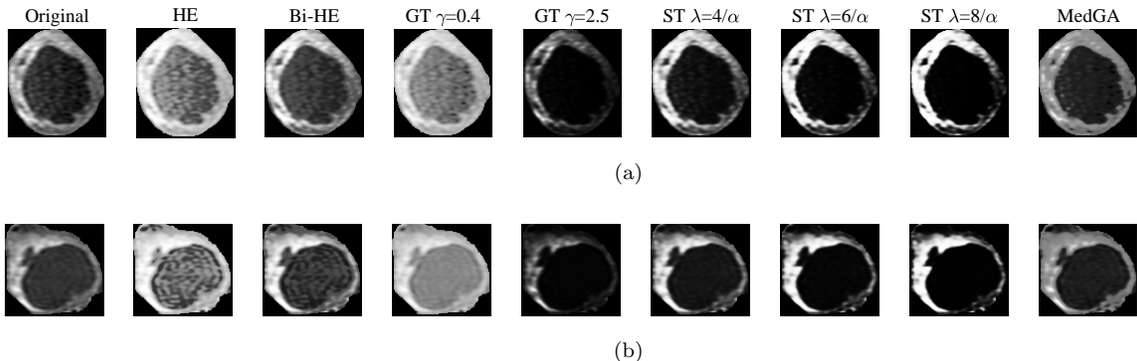


Figure 7: Examples of enhancement achieved on medical images of uterine fibroids by HE, Bi-HE, GT (with $\gamma \in \{0.4, 2.5\}$), ST (with $\lambda \in \{4/\alpha, 6/\alpha, 8/\alpha\}$) and MedGA, compared with the original input MR image.

600 the methods considered in this work. As it can be observed, MedGA strengthens the ROI edges by enhancing details and features useful for image binarization; this result confirms, from a qualitative perspective, the quantitative results presented above.

From an overall view of the metrics, we can claim that the approaches ob-
 605 taining the highest values of the #DE measure (i.e., HE and GT with $\gamma = 2.5$) could imply an over-enhancement of the output image, according to the other image quality metrics. This finding is also confirmed by means of a visual inspection of Fig. 7, where the images enhanced using HE and GT with $\gamma = 2.5$ present an inadequate appearance for image observation and interpretation. To
 610 conclude, MedGA achieves outstanding results in performance evaluation with respect to classic image enhancement techniques.

6. Discussion

The main purpose of state-of-the-art methods for image enhancement is the improvement of the contrast level of the whole image, to obtain a “visually
 615 pleasant” result. On the contrary, MedGA is the first enhancement method that explicitly addresses the challenging issues related to the improvement of medical images that are characterized by a nearly bimodal gray level histogram

distribution. The solution provided by our intelligent image enhancement system can be beneficial to visually assist physicians in interactive decision-making tasks, as well as to improve the final outcome of downstream automated processing pipelines for useful measurements in the clinical practice (Rueckert et al., 2016).

MedGA also deals with the practical problems regarding the interpretability of the results yielded by advanced Machine Learning and Computational Intelligence methods in medicine (Cabitza et al., 2017). Indeed, the final best solution found by MedGA (i.e., the output gray level histogram) and the corresponding enhanced image are understandable by physicians. In addition, the efficient encoding of the individuals—taking inspiration by (Hashemi et al., 2010)—coupled with effective HPC solutions, allows for a clinically feasible computational framework. We designed a specific fitness function to emphasize the two Gaussian distributions composing a bimodal histogram, while the existing approaches based on evolutionary computation or Swarm Intelligence techniques were conceived for a different purpose, i.e., improving the perceived visual information in terms of image contrast. Thanks to this *ad hoc* fitness function—differently to the GP-based image enhancement method in (Poli & Cagnoni, 1997), where no fitness function is defined because the user interactively selects the best solution—and the robustness achieved by means of the calibration step for GA’s parameter setting, MedGA does not require any user interaction step. Moreover, MedGA differs from GP-based approaches, in which the final generated solution could have large size (Castelli et al., 2014), so heavily affecting the readability and interpretability of the provided solutions. The main impact of this contribution consists in showing how an effective method based on evolutionary computation can outperform the existing methods in medical image enhancement.

In this work, we tested and validated our approach on MR images regarding MRgFUS therapy for uterine fibroids. More generally, the application of MedGA can be extended also to real-world problems involving the analysis of images characterized by an underlying bimodal histogram, as in the case of bright-field and fluorescence microscopy imaging (Meijering, 2012).

Considering the achieved results in terms of #DE (see Table 1), MedGA’s
650 performance could be further improved—in terms of contrast—by integrating a
novel component in the fitness function, which explicitly relies on the number of
detected edges. Since this additional component would have a different purpose
and a different magnitude, a multi-objective optimization method should be
taken into account. In particular, MedGA could be extended by means of an
655 effective evolutionary computation approach, such as NSGA-III (Deb & Jain,
2014), to simultaneously optimize both conflicting objectives, which consist in
maximizing the number of edges while minimizing the distance between the
optimal threshold and the two normal distributions.

7. Conclusion and future work

660 A novel image enhancement method based on GAs, specifically tailored for
medical images characterized by a bimodal histogram, was proposed in this
paper. This computational framework, named MedGA, exploits a fitness func-
tion that better reveals the two underlying sub-distributions of the gray level
intensities, consequently allowing for an improvement in the results achieved
665 by threshold-based algorithms. Unlike the traditional image enhancement tech-
niques that generally improve the contrast level of the whole image, MedGA fo-
cuses on MR image sub-regions characterized by a roughly bimodal histogram,
making it valuable in clinical contexts involving CE MRI analysis.

We tested and validated our approach on MR images representing uterine
670 fibroids of patients who underwent MRgFUS therapy. MedGA was compared
against the most common image enhancement techniques, overall achieving the
best performances with respect to both signal and perceived image quality, while
preserving, unlike classic HE techniques, the input mean brightness. This novel
medical image enhancement technique was therefore shown to be a promising
675 solution, suitable for medical expert systems.

We remark that, although MedGA exploits the same encoding of individuals
defined in (Hashemi et al., 2010) and (Draa & Bouaziz, 2014) we did not compare

it to other approaches based on evolutionary computation or Swarm Intelligence techniques (see also (Chen et al., 2018)), since they were conceived for a different
680 purpose, that is, improving the perceived visual information of the whole image. As a matter of fact, these approaches explicitly include in the fitness function both the number of edge pixels and the intensity of such pixels, thus achieving high #DE values that would consistently lead to over-enhanced images.

As a future extension of this work, in the case of large size images (e.g.,
685 1000×1000 pixels) we plan to use Graphics Processing Units, which represent an enabling technology for real-time radiology applications (Eklund et al., 2013), since the running time of histograms computation can be considerably reduced by using a parallel implementation (Scheuermann & Hensley, 2007).

In addition, we plan to integrate MedGA as a pre-processing step within
690 an automatic pipeline defined in the context of MR image classification for efficient computer-assisted segmentation using thresholding techniques, such as (Ridler & Calvard, 1978; Trussell, 1979; Otsu, 1975). Indeed, MR image segmentation is a compelling task in radiology practice, for instance in brain tumor detection and delineation (Sompong & Wongthanavas, 2017). Especially, we
695 plan to apply MedGA to metastatic cancer segmentation in neuro-radiosurgery therapy (Leksell, 1949), wherein the enhancement region must be accurately segmented (Militello et al., 2015a; Rundo et al., 2017). In order to make the resulting expert system fully automatic, the segmentation of the ROI bounding
700 results repeatability during patients follow-up. In the case of a significant data availability, this step may be performed by means of approaches based on Deep Neural Networks (Rajchl et al., 2017). In these clinical scenarios, MedGA can be suitably integrated in expert systems tailored for MRgFUS treatment evaluation (Rundo et al., 2019), and brain tumor segmentation in neuro-radiosurgery
705 (Meier et al., 2016).

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