High-power radiofrequency ablation guided by ablation index for pulmonary vein isolation

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Abstract
Background
Proposed to facilitate pulmonary vein isolation (PVI), high-power ablation may cause extracardiac damage. This study evaluated the safety and efficacy of ablation index (AI) guided high-power ablation first in an animal model and subsequently in a clinical study.

Methods
Outcomes of radiofrequency (RF) applications were compared in a swine ventricular endocardial model (n = 10 each for 50W, 40W and 30W; AI = 500), and in 100 consecutive patients with paroxysmal atrial fibrillation undergoing PVI (40W [last n = 50] vs. 30 W [first n = 50]; target AI = 400/500 on posterior/anterior wall, respectively). Acute PV reconnection was assessed post adenosine administration 20 minutes after ablation.

Results
In swine ventricular endocardial RF applications, use of 50W and 40W vs. 30W was associated with greater tissue lesion depth (5.06 ± 0.16 and 4.38 ± 0.13 mm vs. 3.95 ± 0.16 mm; P < 0.001) and smaller lesion maximum diameter (7.81 ± 0.15 and 8.42 ± 0.18 mm vs. 9.08 ± 0.15 mm; P < 0.001). Tissue necrosis caused by 50W vs. 40W and 30W was the deepest and largest (3.15 ± 0.18 mm vs. 2.71 ± 0.17 and 2.42 ± 0.13 mm; and 5.58 ± 0.18 mm vs. 5.18 ± 0.16 and 3.94 ± 0.17 mm; respectively; P < 0.001). In PVI, use of 40W vs. 30W was associated with shorter procedure time (56.54 ± 1.81 min vs. 76.55 ± 2.34 min; p < 0.001) and ablation time (35.85 ± 14.87 min vs. 51.01 ± 17.99 min; p < 0.001); lower RF energy per point (909.02 ± 354.57J vs. 1045 ± 376.60J; p < 0.001); higher first-pass PVI (87% vs. 72%; P < 0.01); lower acute PV reconnection (22% vs. 41%; P < 0.01); no complications in either group; and similar sinus rhythm maintenance at 12 months (92% vs. 84%; P = 0.22).

Conclusions
AI-guided high-power (40W) vs. conventional (30W) PVI was related to a reduced time for procedure and was considered safe, with diminished acute pulmonary vein reconnection.

Introduction
Ablation through radio frequency (RF) is popularly used in the treating patients with atrial fibrillation (AF) via transmural, continuous, and lasting lesion formation with pulmonary vein isolation (PVI).
Despite optimization of RF current transfer, RF energy transfer duration, stability of catheter and tissue in contact with catheter (1), PV reconnection occurs acutely after PVI, at three months and at 22% and 15% frequencies, respectively (2, 3), mainly secondary to reversible damage, partial thickness and/or incomplete ablation (4–6). Intense power Radio Frequency energy delivery, although, for a shorter length of time, could curtail the negative effects of the instability of catheter, that are inherent, leading to edema in tissue, and possibly enhance the lesion to lesion consistency by altering the link between resistive heating and conductive heating, allowing permanent tissue damage due to immediate heating while reducing the impact of heating due to conduction and as a consequence reduced damage to adjoining structures. To this end, shorter duration 50W lesions have been associated with more effective long-term outcomes with no enhancement in complications rates (7).

Higher intensity ablation constantly resulted in procedures of lesser duration, lesser dose of fluoroscopy, as well as declined delivery of gross Radio Frequency energy Higher-power ablation consistently resulted in procedures of lesser duration, lesser dose of fluoroscopy, as well as declined delivery of total RF energy (8–9). In recent animal studies, 50W ablation for 5 seconds was superior to lower-power, longer-duration ablation for lesion creation with lower complication rates (10, 11). A study conducted recently pointed out that by using contact catheters that are force-sensing and point-by-point ablation, 2-year freedom from paroxysmal and for persistent Atrial Fibrillation after one procedure was 86% and 72%, respectively; per point, the average delivery time of the 50W energy was only 11.2 ± 3.7 seconds, and 895 ± 258 seconds was the total duration of the delivery of Radio Frequency energy (12). Many centers currently perform RF ablation using 25–35W power for 30–60 seconds at each point (13, 14). In the present 2-tiered study, we first tested in a swine model the effect of different power output on lesion creation at the ventricular level and compared lesion dimensions for the same ablation index (AI) value. A clinical study then assessed whether AI guided high-power ablation relative to AI guided standard power ablation is relevant for PVI in cases of paroxysmal atrial fibrillation.

Methods
The present study was divided into 2 successive parts. An animal study provided efficacy and safety
data, which oriented the clinical study for the choice of RF duration. The 2 parts followed the same technical principles for clinical research.

**Animal study**

Swine hearts were obtained from an approved vendor (Shandong Animal Experiment Center) and studied at a laboratory of Yantai Yuhuangding Hospital. After pericardium excision, hearts were kept in a tissue bath with sodium chloride solution circulating at 0.9% at a consistent temperature of 37 °C, and individual radio frequency procedures were conducted at the endocardial aspect of the ventricles. Radio frequency ablation was conducted with a 3.5-mm-tip contact catheter that was force sensing and open irrigated (Thermo-Cool Smart Touch (ST), Biosense Webster, South Diamond Bar, CA). A steady CF, or Contact Force of 10 g was maintained by the operator who proceeded with the energy delivery after positioning the catheter perpendicular to the left ventricular tissue. For the same AI of 500, power output was 30 W for the control lesions and 40 W and 50 W for the study lesions, respectively, at ten ablation points. Ablation sites were >10 mm apart to avoid any lesion overlap and to facilitate histopathologic evaluation. Lesions and necrotic areas were then divided into two perpendicular planes for the calculation of depth and maximum possible internal width. After proper fixation in 10% natural buffered formalin, hearts were trimmed to isolate all ablation sites, which were processed and embedded in paraffin. All paraffin blocks were subjected to microtome dissection twice serially at 5 µm, before being stained with hematoxylin and eosin for histopathologic analysis.

**Clinical study**

**Patient population**

Subjects consisted of symptomatic patients who had consecutive paroxysmal AF undergoing initial point-by-point RF ablation at Yantai Yuhuangding Hospital (Yantai, China) from April 2018 to February 2019. The first 50 patients (control group) were treated with standard power (30 W), whereas the last 50 patients (study group) were treated with high-power (40 W). Study exclusion criteria were left atrial thrombus, ages less than eighteen years, prior Atrial Fibrillation ablation, ejection fraction of less than 35% of the left ventricle, severe coronary artery disease or valvular disease, thyrotoxicosis, and a
greater than 60 mm diameter of left atrium. For the Ablation Procedure, informed and written consent was obtained from all patients. The current study’s protocol was approved by the Yantai Yuhuangding Hospital (Yantai, China) Ethics Committee.

**Ablation protocol**

We have already elaborated the periprocedural anticoagulation protocols earlier(15). The procedure was performed under local anesthesia consisting of administration of lidocaine to the groin and left subclavian region, with the analgesic (fentanyl). The CARTO3 system (Biosense, Webster, Diamond Bar, CA) was used in all cases for 3D mapping. Three catheters were inserted: (1) a 6F decapolar catheter (Biosense Webster, Diamond Bar, CA) was placed into the coronary sinus; (2) a multipolar mapping catheter (Pentaray, Biosense Webster, Diamond Bar, CA) to create atrial anatomy shell; and (3) a 3.5-mm-tipped contact catheter that is force sensing and open irrigated (ThermoCool Smart Touch (ST), Biosense Webster, South Diamond Bar, CA). Following a dual trans-septal puncture, 2 SL1 sheaths were introduced into the left atrium, a sheath for the Pentaray catheter (multipolar mapping catheter) and the other sheath for the Cartographic Information Division (CAETO) ablation catheter. After creating an anatomical CARTO map of the left atrium, the borders of all four pulmonary veins (PVs) were marked on the CARTO map. Next, all 4 PVs underwent encirclement by point-by-point RF applications (Target AI was 400 for the posterior segment, and 500 elsewhere) and other ablations as clinically indicated (Fig. 1). RF applications were conducted with power-control mode, temperature maintained at 45°C, and saline irrigation (17 to 30 mL/min). Ablation was made targeting with a Contact Force of 10-20g. Where Contact Force was regularly less than 5g or greater than 40 g, ablation was paused, and adjustment of catheter position and orientation was undertaken as required. The operator was alerted with a flashing screen visual warning if the Contact Force exceeded 30g. Pulmonary Vein Isolation was the endpoint of the procedure. Blockage of entrance was determined by the lack of Pulmonary Vein potentials recorded with the tissue proximity instruction (TPI) of pentaray catheter (16). Blockage of exit was illustrated as the failure to capture the left atrium while pacing with the TPI of pentaray catheter within the antrum carina included (17). A waiting time of twenty minutes was noticed in both the groups subsequent to the Pulmonary Vein Isolation,
following which an administration of eighteen mg of adenosine was initiated with the Pentaray in place to record Pulmonary Vein Reconnection. In the case of repetitive Pulmonary Vein reconnection, Radio frequency Ablation was performed to ensure isolation. However, in the case of transient reconnection, ablation was made and adenosine was repeatedly administered to ensure that there was no reconnection possibility. The AI Software Module (Biosense Webster Inc) was utilized for identification of Automatic lesion tagging (VISITAG). Ablation was performed with ST catheters in a controlled 40W power mode in the study group and a controlled power mode of 30W in the control group. CARTO was used with the “Distance Ruler” function; Taking the center of the last VISITAG, there is a continuous measurement of distance with changes being recorded in real time (Figure 1). At the commencement of an ablation lesion, the distance was maintained at less than 6 mm. The radius of the VISITAGs were such that 3 mm ablation lesions were displayed and therefore VISITAGs of less than 6 mm could be seen to overlap (Fig. 1). Gaps , if any, between VIDITAGs were looked for subsequent to the completion of the circumferential line of Ablation, making the geometry transparent, indicating that the result would be gaps that are greater than 6 mm. An ablation lesion was used to fill gaps, if any were noticed. The operator’s discretion prevailed whether to redo a lesion, in case there were VISITAGs that were not red (meaning the AI target was not met). In case the pulmonary vein was not isolated after the circumferential line of ablation was completed, more ablation was done under the guidance of the Pentaray catheter either with in the line of ablation or on it where the signal existed. In the intravenous ridge between pulmonary vein, ablation was not conducted, unless Pulmonary Vein Isolation could not be achieved otherwise. First-pass antral isolation designated exit block obtained after initial anatomic encirclement.

**Analysis and Collection of Data**

Gender, CHADS2 and CHA2DS2-VASC scores, age, BMI or Body Mass Index, size of the left atrium, history of transient ischemic attacks or strokes, existence of hypertension, diabetes, coronary artery disease (CAD), cardiomyopathy, and obstructive sleep apnea were recorded for each patient. We calculated the time for procedure per ablation, total RF times, and CF. Reporting of complications was per procedure. Steam pops, phrenic nerve paralysis, strokes occurring from 48 hours to 30 days and
strokes occurring within 48 hours, death, incidence of pericardial tamponade, strokes occurring within 48 hours and, PV stenosis requiring intervention, atrioesophageal fistulas, and catheter char were the complications that were studied. The first ablation point to PVI, not including additional ablation beyond PVI constituted the procedure time. The RF time was defined as the time for which radio frequency energy was used. If there were no atrial fibrillation, flutter, or tachycardia lasting more than 30s off antiarrhythmic drugs after a 3-month blanking period, then that constituted a successful ablation procedure.

**Post procedure Follow-up**

All patients were discharged home within 48 to 72 hours. Vitamin K antagonists or direct oral anticoagulants were prescribed for at least 2 months (subsequent strategy depending on the CHA2DS2-VASc score). At the clinic, patients were given an ECG scan within 24 hours, and then subsequently at the end of three and nine months. Holter monitoring was performed at six months and twelve months and beyond as per the symptoms. Recurrence was defined by any documented episode of AF or atrial tachycardia lasting >30s.

**Statistical analysis**

For each part of the study, the variables were presented as mean ± Standard Deviation or percentages, as deemed fit. Continuous and categorical data were compared with the Student t test (two-tailed) and the χ2 test (or Fisher exact test in case of small sample), respectively. P<0.05 was taken as statistically significant.

**Results**

**Animal Study**

Table 1 shows a comparison of lesions and necrosis in control (30W) and study (40 W, 50W) swine hearts for a total of 30 RF applications. By visual inspection, 50W use was associated with shallower lesions; and increasing RF power and shortening RF duration yielded greater maximum diameter (green line) and larger and deeper tissue necrosis area (red line) and nuclear pyknosis by histopathology (Figure 2).

**Clinical study**
Hundred successive patients with paroxysmal atrial fibrillation were evenly distributed to control group (first 50 patients) and study group (last 50 patients); their baseline clinical characteristics are summarized in Table 2. A total of 8726 Radio frequency lesions were administered to these 100 patients, and procedural data are compiled in Table 3. Figure 3 shows the spread of lesions by ranges of CF (3.74%, <5 g; 86.48%, 5-25g; 2%, >25 g). The study vs. control group had shorter total procedure time (56.54±1.81 vs 76.55±2.34 minutes, p<0.001), left and right encirclement procedure time (25.83±1.12 vs 37.92±1.24 minutes, and 30.92±1.31 vs 38.33±3.06 minutes; both p<0.001), total ablation time required for PVI (35.85±14.87 vs 51.01±17.99 minutes p<0.001), ablation time per point in different parts of PV and total RF energy delivered per procedure (909.02 ± 354.57 vs. 1045 ± 376.60), p<0.001; Figure 4). Figure 5 shows the energy delivery per point throughout PV. Isolation of first pass was more frequent in the study group (87% vs. 72%, p<0.01), occurring in both left and right in 70% PVs (35/50) of study patients vs. 42% (21/50) of controls (p<0.001). The study vs. control group required fewer additional ablations on the intervenous ridge between the pulmonary veins to isolate them (8% [8/100] vs. 21% [21/100], respectively, p<0.001), and had fewer acute PV reconnections (11% [22/200] vs. 23% [46/200], p<0.001).

At mean 1.04±0.62 years follow-up, the percentage of patients free from atrial fibrillation or atrial tachycardia after one procedure was 92% (46/50) in the study group and 84% (42/50) in the control group (p=0.22). There were no cases of PV stenosis, stroke, pericardial tamponade, TIA, atrial-esophageal fistulæ, or death.

Discussion
The present two-tiered animal and clinical study yielded four major findings. First, in swine hearts, higher vs. lower power RF applications were more effective in creating larger and deeper necrosis yet shallower lesions, while lower power was associated with deeper lesions without basophilic changes in connective tissue. Second, in patients with paroxysmal atrial fibrillation, high-power ablation was associated with lesser duration procedures and lesser total radio frequency energy delivery, especially on the left anterior wall. Third, ablation index (AI), the novel marker incorporating time, power and contact force, reliably predicted the degree of necrosis in RF delivery. Fourth, higher power
combined with AI increased PVI effectiveness with more frequent first-pass isolation, decreased acute connection, and favorable 12-month outcomes.

Myocardial lesion creation starts at 45°C, being partially reversible below 50°C (transient stunning), and definitive above 50°C (durable necrosis)(18). Delivery of radio frequency energy to the tissue is a complicated interaction(1). Thermal injury induced by electrical current delivery with an irrigated-tip comprises resistive and the conductive phases. Resistive heating is a relation between the actual current supplied to the tissue, with the resistance of the radio frequency generator which probably occurs relatively early in the radio frequency application. By using lower resistance or greater radio frequency power, better resistive heating can be achieved. For instance, with standard power (25–30W), temperature already rises above 50°C, but tissue necrosis is confined to the first 1 to 1.5 mm from the ablation catheter tip(19). Conductive heating, which is a secondary passive heating of deeper tissue increases with the period of RF applications. An electrically silent scar is obtained when the tissue is heated to 50°C or higher for several seconds thereby obtaining irreversible coagulation necrosis. Force sensing and stability monitoring have considerably facilitated the reproducibility of heat transmission to the tissue(20,21). The balance between power and duration parameters, respectively involved in resistive and conductive heating, therefore has a growing influence on lesion creation. By increasing resistive heating size, high-power (40–50W) may theoretically favor the creation of durable lesions (temperature above 50°C) whose dimensions might be particularly suitable for PVI because antral thickness is consistently below 4 mm(22).

Animal experiments

In the present study in swine left ventricular myocardium, 50W power ablations for short durations at 10g of CF created larger and deeper necrosis than that observed with 40W and 30W of power; however, use of 30W resulted in deeper lesions but without basophilic changes which might render them more susceptible to tissue recovery. Higher power, shorter duration applications were more effective in creating larger and deeper necrosis yet shallower lesions may be due to reduced temperature rise in deeper tissue relative to standard lesions. However, the area of tissue necrosis near the surface of the ablation catheter tip is larger because it is more dependent on resistive
heating, which is proportional to power. The larger diameter of necrosis in study group may lead to complete encirclement of pulmonary veins, due to more effective contiguity with adjacent necrosis. At the same time, it was found that higher power (50W) ablation may lead to more complete cell necrosis. However, nuclear pyknosis was observed only more superficially with low-power ablation, and only cell edema was present in the deep part, which may lead to cell reactivation and consequent acute and late PV reconnection and AF recurrence.

Clinical research

Procedural time

In the present study, high-power ablation was associated with shorter procedural duration due to shorter time required for lesion creation, more first-pass PVI, and fewer acute pulmonary vein reconnections. Nilsson et al(23) reported that ablation with 45W for 20 seconds vs. 30W for 120 seconds was associated with shorter pulmonary vein isolation period, mean fluoroscopy time, radiation dose, and total time of radio frequency application. In ventricles from freshly killed pigs, Goyal et al(24) demonstrated that for 20 g of CF, the time required to generate a 4-mm deep lesion decreased from about 20s at 20W to 6-7 s for 50W. In the present study, under the same pressure, the ablation time per point of the left atrial posterior wall in the high-power group was 8-9s shorter than that in the control group, and 10-13s shorter for the left atrial anterior wall. Because catheter instability in a constantly beating heart may also account for the difficulty to transmit heat to the tissue, RF application time shortening probably optimizes lesion creation by increasing the likelihood of catheter stability throughout the entire RF application, particularly for ablation of the left arterial anterior wall by the appendage ridge on the left artery. The significantly smaller pressure of <10g (60%) used for the left atrial anterior wall than for the other three surfaces helps avoid catheter slippage[Figure 6]. However, low power use requires longer time which increases catheter slippage, and the additional time spent adjusting the catheter may cause discontinuous ablation tissue edema, lower first-pass PVI rate, and increased risk of acute PV reconnections. In the present study, ablation time of the left anterior wall was longer than that of other parts of the left atrium in both study and control groups[Figure 7]; however, it showed lesser time in the study when compared to the control
group likely secondary to shorter time required to adjust the catheter or the ineffective point of ablation (procedure time minus total ablation time).

**AI value**

AI, a novel marker in a weighted formula utilizing time, contact force and power in a weighted formula. The use of contact force targets and ablation output markers such as Force-Time Integral diminishes recurrence and complication rates in cohorts of patients with AF undergoing PVI (3,25,26). Single-center studies on AI-guided ablation have demonstrated a very high first pass PVI rates of 97% to 98% and 2% to 6% pulmonary vein Reconnection rates, which are very low (26,27). In the present study on AI-guided high-power ablation, numbers of first-pass PVI were elevated at 87% albeit lesser when compared to earlier reports. Equivalently, when compared to controls the numbers for acute pulmonary vein reconnection were lower (11%). The targets of AI were arbitrary, with ranges for posterior walls at 380 to 400 while for other areas it was 550. ST catheters delivering high power was the novel combination used in this study. With reference to this, the AI target of 400 for posterior wall and 500 for other places has contributed to higher First Pass PVI rates and lower acute Pulmonary Vein Reconnection rates.

**First-pass pulmonary vein isolation**

In AI guided centered evaluation of High Power Ablation, Dhillon G (28) reported more frequent first-pass PVI. Leshem E (29) compared ablation using 90W for a duration of four seconds to 25W for a duration of twenty seconds. The 90W four second ablation resulted in lesions with full thickness and no gaps in all cases, while the 25W twenty second ablation yielded partially thick lesions with several gaps between the lesions. At 25W for a 20 second duration, “endocardial sparing” seemed to occur due to the catheter tip irrigation, which was perceived to be a scar creation failure since there was not enough resistive heating as a result of cooling due to irrigation, for the tissue to be destroyed. In the present animal study experiment, use of higher power was associated with larger maximum diameter and more thorough tissue necrosis. Greater size and better consistency of the created tissue necrosis may explain why high power increases procedure efficiency by ensuring more first-pass PVI and fewer reconnections at 20 minutes.
Acute pv reconnections

An important requirement for the success of the procedure is the durability of PVI; Late as well as acute pulmonary vein reconnection have been known to cause atrial fibrillation (30,5). In a study using an open irrigated-tip catheter, patients undergoing ablation at 50W vs. 35W had greater freedom from AF (82% vs. 66%)(31). The concept of the “weakest link” indicating locations of reconnection was explained in the recent past by El Haddad and colleague(32), who discovered that the absence of contiguous lesion sets and the insufficiency of lesion depth cause the sites of reconnection. Consistent with other studies, the present one documented fewer acute PV reconnections in the high-vs. standard power group, and in an animal model, higher power caused more thorough tissue necrosis thereby precluding cell restoration and reducing acute pulmonary vein connection.

Complications

By avoiding tissue damage caused by distant conductive heating, which becomes perceptible at later stages of longer duration radio frequency applications, and providing solutions by way of shorter duration, but higher power radio frequency energy delivery, wherein tissue destruction happens at the earlier portion of the Radio Frequency Application . In porcine ventricles of freshly killed pigs, Goyal et al(24) demonstrated that to create a 4 mm deep ablation the time taken was twenty seconds at 20W, whereas at 5W it was a mere 6-7 seconds, thereby implying that collateral injury can very much be reduced by utilizing the higher power lesser duration radio frequency applications approach. Bhaskaran et al(10) demonstrated that a 5 second procedure at 50W achieved transmural lesions which were inherently safer than a 30 second procedure at 40W. While there was a 8% incidence of steam pops in a 30 second procedure at 40W, there were none in a 5 second procedure at 50W. Winkle et al(7) created a comparison between a 3 to 10 second 50W short duration procedure using open tip irrigated catheters and a 25 to 40 second lower power application. There was no increase in complications and there were much greater incidence of freedom from atrial fibrillations long term, with shorter procedural and fluoroscopy times. Winkle(33) compared the incidences of complications at four experienced centers between atrial fibrillation ablations performed at 45W to 50W power intensity over 2 to 15 second durations per lesion, against reduced power 30W procedures over 20
second durations, and concluded that ablations performed at 45W to 50W over short durations yielded results with very low complication rates. Our clinical results cannot provide definitive information (for example using esophageal temperature monitoring) on whether high-power, short-duration lesions were safer than low-power energy delivered for a longer time. However, our histological results confirmed shallower tissue lesions with high-power than low-power ablation, with shorter ablation time reflecting shorter time for catheter to attach to the atrial wall tissue, thus leading to higher safety. In addition, the extremely low complication rate may be reassuring for using 40W in left atrium at short duration, and may encourage considering use of high power short time radio frequency ablations to enjoy the benefit of reduced procedural time, fluoroscopy, and total RF energy delivery times, even on the posterior wall.

**Limitations**

Despite this observational study demonstrating encouraging results, their robustness is limited by several factors. First, degree of esophageal injury during ablation is unclear because all patients were treated under local anesthesia and could not tolerate esophageal temperature monitoring, and no postoperative endoscopic examination was performed. Second, the relatively fixed AI values used may be insufficient or excessive for some patients, especially for thin women. Third, because ST were used instead of SmartTouch Surround Flow (STSF) catheter, there is a small amount of scab at the catheter tip during the 50W ablation in animal study, for which, 50W was not selected for ablation in clinical research. Fourth, after a year, there was similar sinus rhythm maintenance rate after ablation with 30W and 40W which may because of the numbers in each group were probably too small to look at a real difference in these two techniques. However, use of 50W during ablation may further improve long-term prognosis.

**Conclusions**

In the present study of high-power ablation, use of guidance by AI, which provides a rational local endpoint allowing for a tailored short-duration radiofrequency application associated with optimized lesion metrics, translated into shorter procedural time and improved acute efficacy, without compromising safety profile and long-term outcomes relative to standard-power ablation.
Declarations

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Availability of data and materials

The data sets used and/or analyzed during the present study are available from the corresponding author upon reasonable request.

Authors' contributions

ZXF and LJP designed the study. HF, ZXF WCX LJP CHX and ZHH performed the experiments. LWJ and ZHH analysed the data. ZXF and LWJ prepared the manuscript. All authors have seen and approved the final published version of this manuscript.

Ethics approval and consent to participate

All patients enrolled in the study signed informed consent documentation. All experimental procedures were conducted under the approval of the Clinical Experiment Ethics Committee of Yantai Yuhuangding Hospital.

Patient consent for publication

Not applicable.

Competing interests

The authors declare that they have no competing interests.

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Tables

Table 1. Swine Ventricular Lesion And Necrosis Characteristics
| Variable                        | 50w            | 40w            | 30w            | P Value  |
|--------------------------------|----------------|----------------|----------------|----------|
| ablation time per point, s*    | 20.00±1.095    | 27.78±9.72     | 44.83±9.83     | <0.001   |
| Ventricular lesion impedance drop Ω* | 13.5±1.871    | 9.11±1.537     | 6.17±0.983     | <0.001   |
| Energy delivery per point, J*  | 992.33±54.331  | 1085.11±45.018 | 1337.00±24.421 | <0.001   |
| Ventricular tissue lesions depth, mm* | 3.95±0.16     | 4.38±0.13      | 5.06±0.16      | <0.001   |
| Ventricular tissue necrosis depth, mm* | 3.15±0.18     | 2.71±0.17      | 2.42±0.13      | <0.001   |
| Ventricular tissue lesions width, mm* | 9.08±0.15     | 8.42±0.18      | 7.81±0.15      | <0.001   |
| Ventricular tissue necrosis width, mm* | 5.58±0.18     | 5.18±0.16      | 3.94±0.17      | <0.001   |

Values are mean±SD or n (%)
Table 2. Clinical Characteristics

| Variable                        | Study Group; High-Power; (50 Patients) | Control Group; Standard Power; (50 Patients) | P Value |
|---------------------------------|----------------------------------------|---------------------------------------------|---------|
| Age, y*                         | 64.4±9.45                              | 64.9±8.62                                   | 0.862   |
| Male, n (%)                     | 34 (68)                                | 32 (64)                                     | 0.673   |
| LV ejection fraction, %         | 64.74±4.46                             | 61.9±5.40                                   | 0.08    |
| Left atrial size (mm)           | 40.65 ± 5.87                           | 42.85±3.10                                  | 0.147   |
| Hypertension, n (%)             | 22 (44)                                | 26 (52)                                     | 0.423   |
| Diabetes mellitus, n (%)        | 6 (12)                                 | 8 (16)                                      | 0.564   |
| Body mass index                 | 21.27 ± 1.88                           | 21.04 ± 2.65                                | 0.749   |
| Prior stroke/transient ischemic attack, n (%) | 8 (16) | 10 (20) | 0.603 |
| Coronary artery disease, n (%)  | 14 (28)                                | 12 (24)                                     | 0.648   |
| CHADS2 score                    | 2.1 ± 0.64                             | 1.95 ± 0.80                                 | 0.504   |
| CHA2DS2-VASC score              | 3.4 ± 1.14                             | 3.15 ± 1.04                                 | 0.474   |
| Variable                          | Study Group; High Power (40w) 50 Patients | Control Group; Standard Power (30w) 50 Patients | P Value |
|----------------------------------|------------------------------------------|-------------------------------------------------|---------|
| Procedure time, min              | 56.54±1.81                               | 76.55±2.34                                      | <0.001  |
| Right encirclement Procedure time, min | 30.92±1.31                               | 38.33±3.06                                      | <0.001  |
| Left encirclement Procedure time, min | 25.83±1.12                               | 37.92±1.24                                      | <0.001  |
| Right encirclement points n      | 51.25±2.45                               | 49.08±3.58                                      | 0.1     |
| Left encirclement points n       | 44.42±1.44                               | 45.00±1.91                                      | 0.407   |

Values are mean±SD or n (%)
|                                | Group 1 | Group 2 | p-value |
|--------------------------------|---------|---------|---------|
| Total ablation time, min       | 35.85±14.87 | 51.01±17.99 | <0.001 |
| Contact-force, g*              | 12.07±5.34 | 11.85±5.40 | 0.523  |
| Ablation time per point, s     | 22.64±9.39 | 32.56±11.48 | <0.001 |
| Energy delivery perpoint, J*   | 909.02±354.57 | 1045±376.60 | <0.001 |
| Impedance drop per point, Ω*   | 10.13±1.624 | 6.57±1.012 | <0.001 |
| First-pass PVI, n (%)          | 87(87)   | 72(72)   | <0.01  |
| Reconnection after 20 min, n (%)| 22(11)   | 46(23)   | <0.01  |
| Groin hematoma, n .(%)         | 1        | 1        | 1      |
| Tamponade, n (%)               | 0        | 0        | 1      |
| Peri-procedural stroke, n (%)  | 0        | 0        | 1      |
| Esophageal fistula, n (%)      | 0        | 0        | 1      |
| Sinus rhythm at 12 mo, n (%)   | 46 (92)  | 42(84)   | 0.22   |

Values are mean±SD or n (%)
Figure 1

AI-guided ablation injury. VISITAGs placement with Distance Measure Tool.
Figure 2

Ventricular lesions obtained with constant Al of 500 and with 50W, 40W or 30W of power output. A. By visual inspection, increasing power was associated with larger lesion maximum diameter (green line) and deeper tissue necrosis (red line), with boundaries of tissue lesion and necrosis becoming clearer. B-D. Lesions (green line) and necrosis (red line) generated with 30W, 40W and 50W were examined under 20x amplification; under 400x magnification at 1mm from the ablation catheter tip in panels B1 (mainly basophilic changes of connective tissue with least myocardial cell changes and nuclear pyknosis), C1 (small number of myocardial cells with fuzzy sarcoplasm, no horizontal stripes and nuclear pyknosis) and D1 (largest number of affected myocardial cells); and under 200x magnification at 3mm from the ablation catheter tip in panels B2 (no basophilic changes of fibrous connective tissue), C2 (basophilic changes only around cells) and D2 (basophilic changes around the blood vessels and cells).
Figure 3

Percentage of total number of RF lesions by average CF ranges.

Figure 4

Ablation time per point throughout PV.
Figure 5

Energy delivery per point throughout PV.

Figure 6

Distribution of contact-force throughout PV.
Figure 7

Ablation time throughout PV.

Supplementary Files

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