ABSTRACT

Mechanical index (MI) is a measure of acoustic power. It is generally omitted during routine echocardiographic imaging. By adjusting the MI, an echocardiographer can perform various contrast-specific imaging modalities during the same session. (Anatol J Cardiol 2015; 15: 334-6)

Keywords: mechanical index, echocardiography, bioeffect

Introduction

Ultrasound is a type of energy created by pressure waves, and its effects on living tissues are named bioeffects. Heating (thermal) and mechanical (non-thermal) effects are two main bioeffects observed in an ultrasonographic examination. Two indices are derived from these bioeffects in modern ultrasonographic machines, i.e., thermal index (TI) and mechanical index (MI). All ultrasonography machines manufactured after 1992 have to add the two safety indices of MI and TI on their screen.

In this review, we tried to mention about the brief definition and clinical usage of the MI.

Mechanical index

Acoustic power is the amount of acoustic energy per time unit. Acoustic power shows the amplitude of the pulse pressure of the ultrasound beam. The MI gives the operator information about the magnitude of energy administered to a patient during an echocardiographic examination.

The MI is a measure of the power of an ultrasound beam. It is derived from the need for an indicator for the possible non-thermal bioeffects of the acoustic field, such as cavitation and streaming.

Cavitation is the expansion and contraction or collapse of bubbles because of the acoustic pressure of the ultrasound beam. Streaming is the movement of complex fluids because of radiation force pressures. The most important non-thermal bioeffect is cavitation. Therefore, before we introduce the MI, cavitation should be mentioned first.

Microbubbles are small and spherical gas-filled bubbles in the size range of 1-10 microns. When a fluid with a microbubble was exposed to an acoustic field, gas-filled bubbles undergo changes in volume in response to acoustic pressure and get bigger. This phenomenon is known as cavitation. The possibility for cavitation is determined by the frequency of the system. Shorter the interval between pressure waveform, higher is the frequency, and lower is the growth of gas-filled bubbles.

In 1989, Holland et al. showed that the initial size of the cavitation nucleus determined the minimum acoustic pressure waveform amplitude required for significant bubble growth. The initial smaller size of nucleus needs a higher acoustic pressure amplitude to overcome the stronger surface tension. It means that the higher frequencies require a very specific and small bubble radius for cavitation.

The most cavitation sensitive tissues are gas-filled organs such as the lung and intestine.

The MI can be defined as peak negative pressure divided by the square root of the frequency of the ultrasound wave.

It is calculated from a formula: $P_- \div \sqrt{f}$

MI: $P_-$: peak negative pressure (MPascal) $f$: frequency (MHz)

Display of the MI on clinical ultrasonographic machines is mandatory in USA to provide information about the ultrasonographic exposure of the myocardial tissue and to put a limit to the output of the device. However, it is not possible to compare the MI directly from one machine to another; it is one of the most
important setting in a contrast echocardiographic study. At a lower MI, microbubbles start to oscillate and vibrate, thereby giving rise to a signal in ultrasound imaging. In contrast, these microbubbles rupture at a higher MI (4). This microbubble rupture is responsible for the non-thermal bioeffect of ultrasonography (5). The MI displayed on clinical ultrasound scanners is intended to estimate these mechanical bioeffects.

The ideal value of the MI should be <1.9 to avoid the bioeffect. The FDA approved MI value of 1.9 is the maximum threshold for diagnostic imaging. Apfel and Holland (6) showed that bubble growth does not exist below the value of 0.5 for MI. However, cavitation clearly exists for values that are two to three points above this index (6). In the literature, authors reported capillary leakage and extravasations for ultrasound exposures above a MI of 0.4 in vivo tests (7, 8). In a meta-analysis performed by the World Health Organization, ultrasonography did not have adverse fetal effects in routine clinical practice (9). A randomized follow-up trial showed that repeated prenatal ultrasonography had no harmful effects on the fetus (10).

Figure 1 shows parasternal long-axis images at two different acoustic power energies. Echogenicity of the myocardial tissue is brighter at a high MI (1.4) value than a low MI value (0.4). After contrast injection, the scattering of the contrast agent becomes higher as the MI index increases. At high MI values, the contrast agent is destroyed, which causes brighter images. At low MI values, brightness of the contrast agent is low because it is not destroyed rapidly, which results in a longer lasting less brighter contrast effect (Fig. 2).

Role of mechanical index in contrast echocardiography

During contrast echocardiography, agitated saline or second generation contrast agents such as perfluorocarbon are injected intravenously, thereby creating a cloud of microbubbles. Microbubbles are intense ultrasound reflectors and forms a differential reflection compared to surrounding tissues and blood. This differential reflection enables the sonographer to clearly investigate the endocardial border, shunts, and myocardial perfusion. Maximal reflection is dependent on the frequency of the ultrasound beam and the diameter of the microbubble. Standard B mode echocardiography routinely runs at a MI between 0.9 and 1.4, leading to optimal myocardial imaging with variable bubble destruction. The ultrasound beam destroys all perfluorocarbon contrast agents above the MI of 1.3, generating a large amount of acoustic energy and this leads to an instantaneous burst signal effect. A high MI is used to achieve myocardial perfusion imaging. With ongoing destruction, the contrast effect is lost. At lower MIs, particularly <0.4, the ultrasound beam destroys less bubbles. This preserves the contrast effect and
leads to homogenous and continuous imaging of the contrast agents in the blood pool. A low MI is optimal for left ventricular opacification (11).

In a study performed by Holland et al. (12), the authors reported that ultrasonographic contrast agents (UCAs) could significantly lower the threshold required for acoustic cavitation. This finding led to the development of a new idea of using a drug or gene labeled contrast agents for targeting therapeutic applications with much lower ultrasonographic energy. This work suggested that UCAs can be used to induce cavitation for therapeutic applications with much lower ultrasound energy. Many researchers have investigated this field and reported many papers (13-16).

Three methods can be used for drug delivery with UCAs. The first method is to administer microbubbles and the drug at the same time. The second method is to attach the drug to the shell of the microbubbles. The last method is to place the drug into the lumen of the microbubbles.

Drug or gene labeled microbubbles can intentionally rupture with ultrasonographic energy at the target vessel to release the drug or gene to the target site (17, 18). In addition, microbubble destruction by ultrasound wave creates micro jets, forcing drug diffusion to the target tissue (19). Under certain ultrasonographic pressure or MI, microbubbles move and collide with the vascular endothelium. This interaction between microbubbles and endothelium increases the vascular permeability (20). In a study performed by Sorace et al. (21), the authors showed that an increase in the vascular permeability is greatest at a MI of 1.0. Vascular permeability is minimal at a MI of 2.0 because destruction of microbubbles decreases the number of intact microbubbles that interact with vascular endothelium. Therefore, intact and oscillating microbubbles under certain MI are required to increase vascular permeability.

**Conclusion**

As a result, there are no major adverse effects of ultrasonography that have been shown in the use of ultrasonography in routine clinical practice; however, it is advised to pay attention to safety indices such as MI and TI. Therefore, an ultrasonographer or an echocardiographer should be aware of the safety indices of MI and TI. The MI is a measure of ultrasonographic power. By adjusting the power and MI, an echocardiographer can perform various contrast-specific imaging modalities during the same session. By using certain MI values, drug or gene labeled microbubbles can be specifically directed to the target tissue for therapeutic applications.

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