Abstract
Attempts have been made to assess and measure ventricular contractility in patients and whether it can be used to identify heart failure. Due to the assumption that if the contractility of all the muscle fibres in a heart were lower, could it be called heart failure? Early attempts involved the assumption of a model of muscle that had a contractile unit in series with an elastic element, but this was found to be incorrect. Further attempts applied the series elastic model but this model also proved challenging. However, one method has assessed changes in contractility in a given patient, in response to an intervention, but could not compare contractility in a patient with heart failure with a normal person. End-systolic pressure-volume (ESPv) is regarded as a more correct index of contractility and this method was used to confirm changes in contractility from beat to beat during AF; showing results that end-systolic volume varied and indicating a shift of ESPv from beat to beat. This review will discuss the difficulty in measurement, the complicated nature of myocardial fibre orientation and hypertrophy, and whether myocardial contractility failure precipitates increased global heart failure.

Keywords
Heart failure, myocardial contractility, cardiac muscle, end-systolic volume, end-diastolic volume, ejection fraction

Cardiac Muscle Mechanics
The fundamental mechanism of cardiac muscle was identified in strips of heart muscle from which the cell membrane had been removed in a solution where intracellular Ca²⁺ was simulated, so the length-force curve could be measured at different Ca²⁺ concentrations, i.e. complete length-force curves at different levels of contractility via Ca²⁺. The purpose of muscle contraction is to create movement, so one also needed to characterise the way cardiac muscle shortened. Early attempts involved the assumption of a model of muscle that had a contractile unit in series with an elastic element, but this was found to be incorrect and a complete characterisation had to wait for the development of the laser diffraction technique for measuring sarcomere length (SL). Sarcomere shortening velocity has an inverse hyperbolic relationship to force (load), increasing SL increased velocity at any given load except in the absence of any load (maximum velocity Vmax). With an increased contractility (increased Ca²⁺), velocity increased at all levels of load.

An attempt was made to apply the series elastic model to assessing contractility in patients by measuring Vmax as dP/dt/P, where dP/dt is the rate of rise of left ventricular pressure (LVP) and P is the LVP. Not only was the model wrong, but which P do you take? If one takes the developed pressure above diastolic, Vmax starts at zero and dP/dt/P is infinity, which is absurd. If you take the total pressure, dP/dt/P can be anything you like (according to your zero pressure reference value) and Vmax is not of use. A simpler and quite useful index is dP/dt/Pmax, the maximum rate of rise of LVP, which, by chance, is not affected much by end-diastolic volume (EDV) in the intact human, but does reflect changes in contractility. However, one can use this method to assess changes in contractility in a given patient, in response to an intervention, but not compare contractility in a patient with heart failure with a normal person because the rate of pressure rise also depends on synchronicity of contraction of the whole left ventricle.

Frank and Starling
Otto Frank (1865–1944) and Ernest Henry Starling (1866–1927) were contemporary physiologists who both studied the mechanical function of the heart. They both correctly realised that a strip of heart muscle, as with skeletal muscle, developed more force if you increased its length. Starling expressed this as the energy of contraction being dependent on initial length (Starling’s Law), but energy is not the same as force. They also both tried to elicit this property in the whole heart, something that is still a difficult exercise today. Starling did not actually perform the necessary experiment to justify his law, because he showed that stroke volume (SV) increased when one increased the EDV. This is a different property that can be demonstrated in the whole heart, which is nothing to do with force depending on length but shows the heart contracts down to the same end-systolic volume (ESV). Frank first showed that isovolumic pressure increased with increase of EDV, and recently I have been involved in helping a German team who are hoping to publish all Frank’s papers in English. It is already clear from these papers that Frank also measured pressure-volume loops, which were later popularised by Sagawa and Suga.
Pathophysiology

Figure 1: The Difference between the Muscular Force of a Contraction Due to Increase of Length of the Contractile Apparatus (Left) and a Difference in Force Due to a Change in Contractility (Right)

Figure 2: Pressure and Volume at the End of Ejection Are the Same as the Isovolumic Pressure That Can Be Developed at That Volume

While, in theory, one can conceive all pressure-volume loops starting from the same end-diastolic volume (EDV) and varying the pressure at which ejection starts (A), in experimental practice, one has to accept varying EDVs when varying ejection pressure (B). The maximum stiffness of the muscle (elastance) is the tangent to the end-systolic pressure-volume curve (pressure change divided by volume change) and gets less steep with increasing volume.

Measurement of peak isovolumic pressure (Frank curve) was unknown to Starling and is the nearest one can get, in anaesthetised animal experiments, to the force-length relationship of the isolated muscle strip. When the heart is allowed to fill, develop pressure and then eject a stroke volume, one finds the ejection ends at a pressure the same as obtained isovolumically at the same volume.1

I have omitted from this discussion the many papers of Suga and Sagawa because their laboratory always came out with an end-systolic pressure-volume (ESPV) curve that is straight. The slope of the line is called $E_s$, for maximum elastance. The concept of muscular elastance is correct; it expresses the change in force for a given change in length (for the heart the change in pressure for a given change in volume) and is the reciprocal of compliance. Essentially the stiffness (elastance) of the muscle rises and wanes with the time course of the contraction and is proportional to the force. However, I do not think it is linear, so the elastance is the tangent of the pressure-volume curves, which become less steep as the volume increases (Figure 2). Nevertheless, the tangent at any one volume becomes steeper with time during contraction and less steep with time during relaxation.

Pressure-volume loops can be recorded in patients using a catheter in the left ventricle that transduces pressure conventionally but also carries electrodes to measure capacitance as an index of volume. By varying the venous return with a vena caval balloon one can record a series of pressure-volume loops and plot the ESPV. One can also introduce an intervention that increases contractility and record a leftward shift of ESPV. ESPV is regarded as a more correct index of contractility than LVdP/dt max (or maximum acceleration of blood from the left ventricle).6  A reduction in contractility could be induced by getting the subject to breathe carbon dioxide,11 but I am not aware that this has been done while measuring human PV loops. This PV loop method was used to confirm changes in contractility from beat to beat during AF. The difficulty is that ejection pressure is virtually clamped by the aortic pressure, and thus I was not prepared to change LVP with a balloon. Nevertheless, results showed ESV varied, i.e. on strong beats the ventricle emptied more and on weak beats it emptied less, indicating a shift of ESPV from beat to beat.12

Ejection Fraction

Let us imagine the result of a global fall in contractility of the heart, which might be a way of defining heart failure. In Figure 3, this is depicted as a fall to the right of the ESPV. Assuming that fluid retention, if present, has been removed, the heart compensates to keep cardiac output normal by dilating to a new EDV, and thus a reduction in ejection fraction (EF). According to the British Heart Foundation, heart failure is denoted by an EF less than 55 %. Solaro considers that heart failure is a failure of contractility and states, “Disorders of the heart, whatever the cause, remain epidemic in world health, and detailed understanding of control of cardiac contractility is essential in the quest for measures to prevent, diagnose, and treat these disorders.” Solaro13 is that treatments are devised for heart failure that “manipulate...”13 There is vast literature testing various theories about the metabolic substrate that underlies reduced contractility in heart failure.

However, the above example of global fall in contractility of the heart is not realistic. I suppose it might apply to some dilated cardiomyopathies, but what about the common causes, such a coronary disease and the end-stage of untreated hypertension, and in heart failure of hypertrophic cardiomyopathy, I can imagine that contractility is increased. After MI, a scar may form and become a sort of series elastic element for the rest of the heart; therefore, the contractility of the scar is obviously nil, but if you look at the contracting muscle, the contractility may be normal or even increased, maybe due to increased sympato-adrenergic stimulation. Can one regard heart failure, caused by AF, a failure of contractility, when actually, contractility is changing all the time from beat to beat?10

The Confusion of Understanding Heart Failure

A consequence of the adoption of many theories with similar views to Solaro13 is that treatments are devised for heart failure that “manipulate...”
Measuring Contractility in Heart Failure

What Does Heart Failure Do To Cardiac Cells

One can induce heart failure in dogs by pacing the ventricle at 250 BPM for 6 weeks. It has been claimed that myocytes obtained from such animals were longer and thinner than those from controls, and shortening in response to electrical stimulation was reduced.\(^1\) Unfortunately, this cannot be accepted as evidence for reduced contractility in heart failure because: myocytes get damaged in the process of isolation and those from heart-failure animals may be more susceptible to such damage; and shortening is not contractility but load dependent, and the load in these experiments is not known. While in the animal, the load would have been abnormally high due to increased wall force in the dilated heart according to the Laplace equation. In these experiments, the contractility of isolated myocytes is not measurable.

Therapeutic Implications of the ‘Heart Failure Equals Contractility Failure’ Theory

It was logical to counteract supposed contractility failure with a drug that increases contractility (‘positively inotropic’). Such a drug should improve contraction whether or not there is contractility failure. Adrenergic stimulation also increases heart rate and both effects increase the demand of the heart for oxygen, so these drugs should only be used for testing, e.g. dobutamine during echocardiography if one is looking for poorly contracting segments that might respond. However, one can also increase contractility with digoxin and with the phosphodiesterase 3 inhibitors, of which the best known is milrinone. Digoxin (which Frank found to be positively inotropic)\(^2\) is very useful in controlling heart rate stimulation also increases heart rate and both effects increase the demand of the heart for oxygen, so these drugs should only be used for testing, e.g. dobutamine during echocardiography if one is looking for poorly contracting segments that might respond. However, one can also increase contractility with digoxin and with the phosphodiesterase 3 inhibitors, of which the best known is milrinone. Digoxin (which Frank found to be positively inotropic)\(^2\) is very useful in controlling heart rate.

Milrinone has the advantage that it is effective in the presence of beta-blockade with no effect on heart rate.\(^2\) Digoxin and milrinone should probably only be used in acute heart failure as a temporary measure, because they work by increasing intracellular Ca\(^{2+}\). This, in turn, increases the risk of arrhythmias and when given chronically increases mortality, giving milrinone the nickname ‘killrinone’. There are also reports of poor outcomes with digoxin given during sinus rhythm.\(^3\) However, improved mortality rate is not everything and in a disease that is fatal, a drug like this might improve the quality of a shorter life. Another approach is to keep intracellular Ca\(^{2+}\) constant but increase the sensitivity of the contractile proteins to Ca\(^{2+}\), thus predicting a lesser risk of arrhythmias.\(^4\) A similar drug produced a benefit in dogs with tachycardia-induced heart failure,\(^5\) but I am not aware of its clinical development, possibly because it induced arrhythmias in rat hearts.\(^6\)

Conclusion

Possibly, heart failure in dilated cardiomyopathies is due to a global reduction in ventricular contractility. In other types of heart failure, the presence or absence of reduced contractility is unknown. In particular, if the failing heart is subjected to coronary disease in some parts but not others, it is very difficult to determine the contractile state of the unaffected parts. In the case of anatomical abnormalities, e.g. valvular disease and congenital heart disease, the heart may be failing globally but myocardial contractility may be normal. Maybe myocardial contractility failure precipitates increased global failure? With the difficulty in measurement, the complicated nature of myocardial fibre orientation and hypertrophy, it is impossible for me to give a firm opinion. Another paper could be written reviewing all that has been written on these aspects, and are still being written.

References

1. Kentish JC, ter Keurs HE, Ricciardi L, et al. Comparison between the sarcomere length-force relations of intact and skinned trabeculae from rat right ventricle: influence of calcium concentrations on these relations. Circ Res 1984; 55:755-68. DOI: 10.1161/01.RES.55.5.755; PMID: 371928

2. Noble M, Else W. Re-examination of the applicability of the Hill model of muscle to cat myocardium. Circ Res 1972;31:580-9. DOI: 10.1161/01.RES.31.4.580; PMID: 5075377

3. Danielis M, Noble M, ter Keurs HE, Wolfratt B. Velocity of sarcomere shortening in rat cardiac muscle: relationship of force, sarcomere length, calcium and time. J Physiol 1984;395:867-81. DOI: 10.1113/jphysiol.1984.sp015420; PMID: 6499996

4. Noble M. Problems concerning the application of concepts of muscle mechanics to the determination of the contractile state of the heart. Circulation 1972;45:252-6. DOI: 10.1161/01.CIR.45.2.252; PMID: 5000471

5. Drake-Olaid A, Mills CJ, Noble MI, Pugh S. Response
to changes in filling and contractility of indices of human left ventricular mechanical performance. | Physiol 1990;422:29–39. DOI: 10.1153/physiol.1990.sp017970; PMID: 1972191.

6. Frank O. [On the dynamics of cardiac muscle.] Zeitschrift für Biologie 1897;32:370–447 [in German].

7. Starling EH. Lecture on the law of the heart. London: Longmans, Green and Co, 1916.

8. Sagawa K, Saga H, Shoukas AA, Bakalar KM. End-systolic pressure/volume ratio: a new index of ventricular contractility. Am J Cardio 1977;40(5):748–53.

9. Weber KT, Janicki JS, Hefner LL. Left ventricular force-length relations of isovolumic and ejecting contractions. Am J Physiol 1976;231:337–43. PMID: 961884.

10. Bennett ED, Else W, Miller GA, et al. Maximum acceleration of blood from the left ventricle of patients with ischaemic heart disease. Clin Sci Mol Med 1974;46:49–59. DOI: 10.1042/cs0460049; PMID: 4811873.

11. van den Bos GC, Drake AJ, Noble MI. The effect of carbon dioxide upon myocardial contractile performance blood flow and oxygen consumption. | Physiol 1979;287:149–61. DOI: 10.1113/physiol.1979.sp012651; PMID: 430387.

12. Brookes CI, White PA, Staples M, et al. Myocardial contractility is not constant during spontaneous atrial fibrillation in patients. Circulation 1998;98:1762–8. DOI: 10.1161/10166.1154-11615; PMID: 9788821.

13. Solero R. Heart failure as a failure of contractility In: Solaro RJ. Regulation of Cardiac Contractility. San Rafael, CA: Morgan & Claypool Life Sciences, 2011.

14. Dorn GW, Molkentin JD. Manipulating cardiac contractility in heart failure: data from mice to men. Circulation 2004;109:150–8. DOI: 10.1161/01.CIR.0000111581.15521.F5; PMID: 14736503.

15. Hall SA, Cigarroa CG, Marcoux L, et al. Time course of improvement in left ventricular function, mass and geometry in patients with congestive heart failure treated with beta-adrenergic blockade. | Am Coll Cardiol 1995;25:1154–61. DOI: 10.1016/0735-1097(94)00543-Y; PMID: 7897129.

16. Solaro RJ: Heart failure as a failure of contractility. In: Solaro RJ. Regulation of Cardiac Contractility. San Rafael, CA: Morgan & Claypool Life Sciences, 2011.

17. Ravens U, Davia K, Davies CH, et al. Tachycardia-induced failure alters contractile properties of canine ventricular myocytes. Cardiovasc Res 1996;32:13–21. DOI: 10.1016/S0008-6363(96)00121-6; PMID: 8891522.

18. Frank O. Die Wirkung von Digitalis (Helleborein) auf das Herz. Sitzungsber. Gesell Morph Physiol 1898;14:14–43.

19. de Graeff PA, Reuter M, Hell K. Value of digoxin in heart failure and sinus rhythm: new features of an old drug? | Am Coll Cardiol 1996;24:213–9. DOI: 10.1016/0735-1097(96)00247-1; PMID: 8837553.

20. Li KI, Chen H, Zhang J, et al. Preliminary results of a study of the effects of milrinone infusion in the presence of beta-adrenergic blockade. Abstracts of the International Congress & Symposium Series 1991;172:31–34.

21. Lee JA, Allen DG. EMD 53998 sensitizes the contractile proteins to calcium in intact ferret ventricular muscle. | Circ 1997;99:297–306. DOI: 10.1161/01.RES.99.4.297; PMID: 9336485.

22. Drake-Holland AJ, Lee JA, Hynd J, et al. Beneficial effect of the calcium-sensitizing drug EMD 57033 in a canine model of dilated heart failure. | Circ 1997;99:215–18. DOI: 10.1042/circ990213; PMID: 9336485.

23. Evans JS, Lee JA, Jones JR. EMD 57033 enhances arrhythmias associated with increased wall stress in the working rat heart. | Circ 1995;91:59–67. DOI: 10.1042/ cir95059; PMID: 7571569.