How does the Extracellular Matrix Change in the Setting of Heart Failure?

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Abstract
The Extracellular Matrix is a dynamic entity, showing constant degradation and deposition while providing the framework for the cardiomyocytes and interstitial proteins to lie on. Its function is important for the proper myocyte alignment within the heart and for internal communication from cell to matrix. Disregulation of the remodeling process resulting in the breakdown of collagen by matrix metalloproteinases is a hallmark of heart failure pathophysiology and produces functional changes encompassing all matrix proteins. Several etiologies with distinct mechanisms ultimately bring about signs of heart exhaustion such as reduced ejection fraction, reduced compliance and ventricular dilatation. Discussed in this paper is the role of inflammation, collagen cross-linking and of myofibroblasts in matrix dysfunction and the mechanisms with which these changes occur in heart failure. Understanding extracellular protein roles within this context would allow for specific drug targeting and thus prevention of heart failure in the early stages of the disease. More studies must be conducted to discover the specific matrix proteins and cytokines that modulate the pathological remodeling process. Serum biomarkers of extracellular degradation products, selective metalloproteinase inhibitors and a personalized treatment approach with a revival of the current classification of heart failure are topics requiring further exploration.

Key Words: Heart; Heart Failure; Extracellular Matrix; Cardiomyocytes; Matrix Metalloproteinases (Source: MeSH-NLM).

Introduction
Heart Failure is a global problem that is on the rise in the developed world. Worldwide, 38 million people are living with this disease, with a 50% increase within the last 15 years alone.¹ With medical advances improving outcomes from myocardial infarctions (MI) and congenital cardiac conditions, the same advances have also inadvertently brought about a steep increase in the prevalence of chronic heart failure.² For the elderly in England and Wales prognosis can be worse than most cancers, with a 10% 30-day mortality following admission, highlighting the need to improve our understanding and treatment of this progressive disease.³ The extracellular matrix (ECM) is a scaffold including extracellular proteins, in which cardiac cells and cardiac vessels are arranged. We are now aware that the ECM is not a static structure but is actively broken down and reformed by an array of chemical and physical signaling within the heart.⁴ Changes in the myocardial matrix even encompass the cardiomyocytes, the resident cells of the myocardium that respond with a state of pathological hypertrophy.⁵ With heart failure, we see changes to the ECM that may differ in etiology, but all progressively results in an impaired contractility, fibrosis and eventual cavity dilatation.⁶ This review describes the ECM environment, identifies key modulators of its remodeling process in heart disease and finally mentions the clinical implications and future directions of this ever-growing field.

The Extracellular Environment
The ECM composition is vital for the heart’s structure as it regulates the alignment and activity of the cardiomyocytes, which are the resident cardiac contractility, which require a stable interstitium to properly function.⁷ Fibroblasts make up over 70% of cells in the myocardium and functionally secretes the most matrix proteins which is suggestive of the large quantities constantly being made in the interstitial space.⁸ The tight space between the myocytes must be able to transfer signals from the matrix to cell and also convert the individual contracting myofibrils into a collective pump, via the Collagen-Integrin-Cytoskeleton-Myofibril relationship.⁹ The ECM includes structural, adhesive and regulatory proteins that work together to maintain the heart’s normal function.¹⁰ It is vital that all these proteins within the ECM stay balanced as they hold important roles in the maintenance of tissue homeostasis.

Structural Proteins
Fibrillar Collagen: In the heart, we can find collagen types I, III, IV, V, VI and VIII of which types I and III dominate the matrix. Type I provides tensile strength and type III distensibility:¹¹¹² I, II, III, V and VI all have a common fibrillar structure and are not found on the basement membrane of cells.¹³ They all contain one main triple helix chain of Glycine-X-Y repeats where X is usually proline and Y can be any amino acid. The collagen helix terminates into 2 globular side chains on either side, referred to as the N- and C-Propeptides and these help in the alignment of the collagen strands as they are secreted from the cell.¹³ Upon maturation, the side-chains are cleaved by procollagen peptidases, releasing the Propeptides into the ECM and rendering the mature collagen insoluble.¹⁴ Cross-links between collagen fibrils are formed via the copper-containing enzyme, lysyl oxidase.¹⁵ This is a physiological occurrence that helps protect the collagen fibrils, both against the mechanical strains imposed on them as a result of the heart’s dynamic function, but also to enzymatic proteolysis resulting in degradation. Figure 1 shows the structure and cleavage site of a procollagen molecule. Collagen that has formed these cross-links is extremely stable and, pound for pound, is stronger than steel.¹⁶ The collagen in the ECM is constantly being remodeled under the control of fibroblasts that communicate with this scaffold via Discoidin Domain Receptor 2 (DDR2) surface receptors. These receptors allow collagen synthesis and degradation to be finely monitored by fibroblasts.¹⁷ The correct concentration of fibrillar collagen is vital for the heart’s form and function, and an inappropriate amount of deposition is associated

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Editor: Milena-Alexandru Goman
Submission: Aug 8, 2018
Acceptance: Dec 14, 2018
Publication: Dec 22, 2018
Process: Peer-reviewed
with cardiovascular disease and more specifically diastolic heart failure.\textsuperscript{18}

\textbf{Elastin}: Elastin, unlike collagen, is not subject to a fixed turnover rate, and in a healthy myocardium can persist life-long.\textsuperscript{19} Elastin is implanted within its own scaffold of glycoprotein microfibrils once synthesized by fibroblasts, cardiomyocytes or endothelial cells.\textsuperscript{20} The stiffness of the collagen fibers is counteracted upon by the elastic recoil properties of elastin, being up to a thousand times more flexible than collagen. This is what gives the heart the ability to rebound in diastole following a contraction. Elastin is also vulnerable to degradation in cardiac disease, either due to chronic stretch-recoil or due to increased protease activity of Matrix Metalloproteinases (MMPs).\textsuperscript{21}

\textbf{Figure 1.} Formation of Mature Collagen from its Pre-Collagen Precursor, Releasing the Pro-Peptides at the N- and C-Terminals and in this Way Becoming more Stable and Resistant to Degradation.

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{figure1.png}
\end{figure}

\textbf{Adhesive Proteins}

These include all the non-collagenous proteins and account for over 95\% of all the molecules in the ECM.\textsuperscript{22} They are vital for the communication between cells and between the cell and the matrix.\textsuperscript{25} Yonggang et al studied their function in the setting of cardiac injury and concluded that they have a part to play in the response to injury, but individual roles and functions have yet to be identified.\textsuperscript{24} These proteins allow cardiomyocytes to monitor the myocardium and therefore modulate their behavior accordingly, through complex feedback loops.\textsuperscript{25}

\textbf{Collagen IV}: This type of collagen is only found outside the membrane of cardiomyocytes where it folds into a sheet across the basal lamina.\textsuperscript{26} Collagen IV connects the contractile myofibril within the cardiomyocytes with the ECM and is therefore vital for the transfer of force across the cardiac tissue.\textsuperscript{27}

\textbf{Integrin}: This cell-interstitial linking protein is made up of an \(\alpha\) and \(\beta\) transmembrane subunit, as shown in Figure 2, and is required for nearly all communication within the ECM.\textsuperscript{28} Ross and Borge’s review on integrin role and function concludes that they are not merely adhesive proteins but have an important role in cell transduction in health and disease.\textsuperscript{29} Shed integrins have been found in heart failure by Ding et al, cleaved from distressed cardiomyocytes and dysregulating communication within the heart.\textsuperscript{30} Targeting this molecule for treatment has not been successful as there is complex sub-specialization within the integrin family where certain heterodimers bind fibronectin (\(\alpha\beta\)) while others bind to collagen and laminin (\(\alpha\beta\beta\)) and fibronectin.\textsuperscript{31}

\textbf{Fibronectin}: This molecule exists as 3 different subtypes and can exist intracellularly and within the ECM (seen in Figure 2). It has been shown by Sharma et al to unfold itself and thus bind to integrin from within the cell’s basal lamina. In this way it assists in cell migration and differentiation. It is also produced following an injury by fibroblasts, macrophages and endothelial cells to help link up the cells that form the scaffold in the injured area.\textsuperscript{22}

\textbf{SPARC (Osteonectin)}: SPARC stands for secreted protein acidic and rich in cysteine and is found in the basal lamina with many important functions. It stabilizes the collagen matrix by phosphorylation of the SMAD2 pathway with which it controls procollagen secretion out of the cell.\textsuperscript{32} It also facilitates cytokine and growth-hormone efficacy, regulates MMP expression and can alter cell shape.\textsuperscript{33,34} SPARC levels will markedly increase following cardiac injury, acute and chronic.\textsuperscript{34}

\textbf{Figure 2.} The Complex Cytoskeletal Matrix Interaction as well as the Role of the Fibroblast in Contact with the Collagen Matrix via Discoidin Domain Receptor 2 Receptors and with the Myocyte via Integrins.\textsuperscript{24,35} DDR2 is the Discoidin Domain Receptor 2.

\begin{figure}[h]
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\includegraphics[width=\textwidth]{figure2.png}
\end{figure}

\textbf{Collagen Remodeling: Synthesis and Degradation}

One of the most noticeable changes to the ECM in the setting of heart failure is the dysregulation of collagen deposition and degradation. This fine control is performed by Matrix Metalloproteinases (MMPs) and Tissue Inhibitors of Metalloproteinases (TIMPs), two enzymes both formed within the fibroblast.\textsuperscript{36} The former is a protease enzyme and the latter its endogenous inhibitor with both being controlled by fibroblasts within the ECM. MMPs and TIMPs keep a homeostatic control over the collagenous matrix of the ECM as well as over the rest of the matrix proteins. As a result, the ECM is a dynamic entity constantly being broken down and reformed, in both health and disease. What separates a physiological state from a pathological one is the balance of the same remodeling process.

\textbf{Matrix Metalloproteinases}

The matrix metalloproteinases are a family of over 24 zinc-containing endopeptidase enzymes. They are present in cardiovascular disease but can also be found in fibrosis and inflammation all over the body, as well as in tumor metastases.\textsuperscript{37} There are 6 main subtypes of MMP, and their enzymatic action dictates their title. They are all mentioned in Table 1 along with the substrate they degrade and the effects they produce. MMPs are released into the ECM as inactivezymogen molecules where they are activated by proteases such as plasmin or by another MMP.\textsuperscript{38,39} Spinale states that the catalytic domain is similar across all subtypes, but the extracellular binding domain, the C-terminal, is specific and conveys to each MMP its independent properties.\textsuperscript{40} In Table 1 are shown MMP-2 and MMP-9, the Gelatinases, but can even break down Collagen types I, IV, V and the contractile apparatus from within a cell.\textsuperscript{41} Another MMP of importance in heart failure is MMP-14 or otherwise Membrane type-1 MMP (MT1-MMP). This class of protease, unlike the rest, exists only in an active form and can degrade all basement proteins as well as collagen.\textsuperscript{42} Yagi as well as Nagase have both written an extensive list on the cytokines and hormones that they believe activate each MMP and have linked increased MMP activity to a heightened risk of heart failure (shown in...
Table 1). A factor that must be considered is the complex interaction between the ECM components and how they are all intertwined. For example, it has been shown that MMPs might produce a shielding effect to disease, through an unclear downstream effect. Some MMPs, such as MMP2, have shown pro-fibrotic capabilities and may actually cause a net increase in collagen deposition rather than what we would expect, most likely via the TGF-β pathway. To conclude, MMPs perform a diverse set of functions. On one hand they are able to degrade collagen but on the other hand also function to activate fibroblasts and increase collagen degradation. Radauceau et al have discovered that ECM turnover is an independent predictor of all-cause mortality and heart failure hospitalization, therefore a very important aspect in the pathophysiology of Heart Failure. More studies into this field are necessary to more accurately elucidate individual MMP roles.

Tissue Inhibitor of Metalloproteinases

The TIMPs are folded disulphide-linked proteins that contain 2 domains: a C-terminal and an active N-terminal which binds to the active site of MMPs. This binding occurs due to cysteine residues at the N-terminal that react with the Zinc ion within the MMP active site. TIMP inhibition of MMPs has been shown to over-power MMP activity as certain studies of left ventricular hypertrophy and chronic heart failure where both MMP and TIMP levels were similarly increased resulted in fibrotic deposition and not degradation. Perhaps this is a marker of TIMP efficacy being superior in the balance of MMPs and TIMPS. TIMP-4 is believed to be a key player in terms of fibrotic deposition in the heart by Li et al, Schultz et al and Graham et al. Moore states that TIMP-2 and TIMP-3 are most abundant while TIMP-4 is not cardiospecific and only partly expressed by healthy cardiomyocytes. These minor discrepancies in the results could reflect the fact that more studies are required in order to understand the exact roles and populations of these protease inhibitors.

ECM Turnover in Heart Failure

In a failing heart, many pathological events are occurring that have a large impact on the constituents of the heart's tissues. Both the ECM and the cells of the heart are affected by an increase in inflammatory mediators and an increased dynamic load. We now know fibroblasts have the potential to signal pro-inflammatory mediators in the absence of infection or ischemia. This signaling occurs as a result of the heart's inability to cope with mechanical demands and directly impacts the cardiac molecular environment. It is now well documented that ECM remodeling is a distinguished feature of cardiac risk that will gradually lead to heart failure. The key players we tend to focus on are the MMPs that have a direct functional role in the progression to heart failure by enzymatically breaking down the structure of the collagenous matrix. The question that remains is why do we see fibrotic deposition in some cases while in others we see collagen degradation? An interpretation of these seemingly conflicting phenotypes is in line with Kim et al's transgenic mouse model. Mice given an active MMP-1 gene initially responded with myocyte hypertrophy, myofibroblast formation and increased collagen synthesis at 6-weeks. This matches the initial state of hypertrophy and fibrosis seen in early stage heart failure. However, at 12-months the collagen content of the transgenic mice was far lower than the wild-type control mice. The proposed mechanism is one of adapting to the stressor, this case the collagenase MMP-1, by synthesizing more collagen to the point of exhaustion. At this point we begin to see a destruction of the matrix and weakening of the heart wall, markers that we also see in human patients suffering from heart failure. These matrix changes directly influence the heart's mechanical functioning and lead to the gradual signs and symptoms we associate with cardiac failure.

Inflammation mediates ECM Remodeling

As we know, the ECM is constantly active and subject to remodeling and as a result is exceptionally susceptible to any chemical or mechanical stressors that might affect it. Inflammation due to increased cardiac stress induces cytokines such as TGF-β, IL-1β, FGF, NO, and TNF-α, which upregulate MMP activity and increase the migration of myofibroblasts into the heart. Damage associated molecular patterns (DAMPs) released from distressed cells promote further inflammation and were shown to activate fibroblasts in vitro. Bradham et al also found that myofibroblastic cardiac tissue and discovered decreased MMP activity in tissue injected with a TNF-α blocker. This suggests that TNF-α and potentially other cytokines and hormones do interact with ECM turnover and can worsen

| Table 1. Description of the Target and Effect of Each of the Main Matrix Metalloproteinases Implicated in Heart Failure. |
| | MMP | Enzyme Family | Substrate | Result of Activation |
| MMP-1 | Collagenase | C-Terminal Collagen | ↑ Collagen breakdown |
| | | Basement Membrane Proteins | ↑ Myocardial Fibrosis |
| | | | Pro-Inflammatory |
| | | | Mesenchymal Cell Differentiation |
| MMP-2 | Gelatinase | Chondroitin Sulphate Proteoglycan | Angiostatin Cell Apoptosis |
| | | | Collagen III | ↑ TGF-β |
| | | | | ↑ Risk of Heart Failure | ↑ IG-1 |
| MMP-3 | Stromelysin | Basement Membrane Proteins | Epithelial cell Apoptosis |
| | | | Plasminogen | ↑ IG-1 |
| | | | Ecadherin | Pro-Inflammatory |
| | | | Perlecain | Anti-Inflammatory |
| MMP-7 | Matrilysin | Fas Ligand | Adipocyte Differentiation |
| | | | Rank Ligand | Fas-mediated Apoptosis |
| | | | ECM proteins | Pro-Inflammatory |
| | | | Precursor of TNF-α | Chemotactrant |
| MMP-9 | Gelatinase | Precursor of TGF-β | ↑ TGF-β |
| | | Citrate Synthase | ↓ IL-2 |
| | | ECM proteins | ↓ Mitochondrial Function |
| | | Collagen IV | Pro-Inflammatory |
| | | | Plasminogen | ↑ Fibroblast migration |
| MMP-12 | Macrophage Elastase | ECM proteins | Pro-Inflammatory |
| | | Plasminogen | ↑ Fibroblast migration |
| | | Elastin | Endostatin-like fragment |
| MT1-MMP (MMP-14) | Stromelysin | ECM proteins | ↑ Myocardial Fibrosis |
| | | Collagen I | ↑ Risk of Heart Failure |

Legend: ↑ IG-1 means an increase in IG-1 as a result of that MMP being activated. ECM Proteins accounts for osteopontin, fibronectin, hyaluronan, tenasin-C and galectin-9 in these studies.

Angiotensin II and Endothelin-1, which upregulate MMP activity and increase the migration of myofibroblasts into the heart. Damage associated molecular patterns (DAMPs) released from distressed cells promote further inflammation and were shown to activate fibroblasts in vitro. Bradham et al also found that myofibroblastic cardiac tissue and discovered decreased MMP activity in tissue injected with a TNF-α blocker. This suggests that TNF-α and potentially other cytokines and hormones do interact with ECM turnover and can worsen.
Review

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the progression to heart failure by directly interfering with the MMP/TIMP control of the ECM. It should be stated that some attempts to tackle these mediators are being pursued in current specialized centres.22 This inflammatory response is not limited to collagen but also affects fibronectin, osteonectin, elastin and laminin, which were all found to be increased in heart failure pathology by Hein et al.58 Finally, inflammatory mediators can serve as transcription factor activators for MMPs, tipping the scale of collagen homeostasis by direct MMP upregulation.64

The role of the Myofibroblast

Myofibroblasts, which are commonly absent in healthy myocardium having been identified only in heart valve leaflets, are key drivers of ECM remodeling in disease (Progenitor populations shown in Figure 3).66,67 They perform all the functions of an active fibroblast to an even greater degree to cause and increase in matrix deposition and fibrosis.65 They also exhibit a myocyte-like phenotype with contractile capabilities and the generation of a stress-fiber network.67 The matrix deposited by these cells is also said to differ to that of fibroblast origin, with particularly increased amounts of collagen I and fibronectin.68 All these features lead to a change in the normal environment both structurally and composition wise in the setting of heart failure. In the acute setting, myofibroblasts are generated from cardiac pericytes, fibroblasts and smooth muscle cells. Long-term stressor persistence allows myofibroblasts to form from cardiac endothelial cells that transition into mesenchymal cells and also from cells derived from the hematopoietic stem cell line (Figure 3).69 Myofibroblasts contribute to pathological hypertrophy, usually with accompanied diastolic dysfunction.65 This dysfunction is a direct result of the fibrotic stiffness brought about by the myofibroblasts, as described above. Overall, the changes described result in a stiffer, less compliant myocardium that is less effective in its ability to pump blood around the body.70 Myofibroblasts damage the cardiac environment in heart failure but in an MI can have beneficial properties and be necessary for wound healing.24 Interestingly, several animal studies have been pursued in the beneficial treatment using myofibroblast differentiation from progenitor populations such as BMDCs in the response to acute ischemic damage post-MI.71

Collagen Cross-linking in Heart Failure

Collagen Cross-linking (CCL) is a feature of ECM remodeling in heart failure.65,68 and has been shown by Graham et al to be more indicative of a worsening prognosis than collagen type ratios in the ECM.70 CCL occurs via two different mechanisms in the heart: an enzymemediated pathway and a pathologial proteoglycan pathway.72 The latter is most evident with aging and in diabetics, but can occur incidentally with other groups too.73 The end result is an increased stiffness of the myocardium and ventricular dilatation, both increasing the risk of heart failure.74 As previously described, collagen must be cross-linked to increase its strength and resistance to degradation, and a collagen matrix with decreased CCL is a weaker and more unstable structure. Gunja-Smith et al investigated the amount of pyridinoline CCL in human heart collagen directly post cadaveric transplant and found that even though collagen levels were quadrupled in some specimen, the overall CCL between that collagen was poor, resulting in dilatation and decreased compliance, both clinical features of heart disease.75 This study is suggestive of the fact that even though a fibrotic state exists in heart failure (study in question looked at dilated cardiomyopathy), the collagen of these patients although plentiful is weak as a result of its poor cross-linking.

Future Directions of ECM Remodeling Research

ECM Plasma Profiling as a prognostic measure of Heart Failure

In current specialized centers, it is standard practice for multiple cardiac biomarkers to be monitored in patients with heart failure. Several biomarkers including BNP, NTpro BNP, Tropinins as well as proteins like Galectin-3 are screened for to help in disease progression.89-91

Figure 3. The Different Progenitor Populations that Differentiate into Myofibroblasts in the Setting of Injury and Mediate further Remodeling.36 35 36

<table>
<thead>
<tr>
<th>MFT</th>
<th>BMDCs</th>
<th>EMT</th>
</tr>
</thead>
<tbody>
<tr>
<td>Monocyte</td>
<td>Fibrocyte &amp; MSCs</td>
<td>Endothelial Cell</td>
</tr>
</tbody>
</table>

Legend: MFT: Monocyte Fibroblast Transition, BMDC: Bone Marrow Derived Cell, EMT: Endothelial-Mesenchymal Transition, MSC: Mesenchymal Stem Cell.

With the recent advances of cardiac matrix proteomics, ECM biomarkers may also be monitored. As previously described, collagen Propeptides are released into the ECM and are picked up in blood tests as a measure of collagen synthesis. Collagen telopeptides on the other hand should not be cleaved in the physiological state, therefore C and N-telopeptides in serum can point at pathological collagen degradation.82 These biomarkers are good indicators of what is going on in the ECM and are found in our bloodstream before any physical symptom of heart failure is apparent. Using these biomarkers in a day-to-day clinic for cardiac observation should be explored further as the Nterminal propeptide of Type II and the Cterminal of Type III Collagen have been shown to statistically correlate with myocardial fibrosis.65,84,85 Furthermore, Kato et al highlighted the prognostic potential of using other ECM biomarkers in the clinic as osteopontin and MMP/TIMP ratios all decrease significantly post-LVAD (Left Ventricular Assist Device) implantation in patients with heart failure and were found to be consistently higher in those who suffered with right ventricular failure in the months following the surgery.85 A final reason to further explore this area of research, as opposed to looking at the ECM under a microscope, are the barriers with attaining live cardiac biopsies which although proven relatively safe in practice, having them performed for research purposes is not ethically justifiable.86 A Review written by Mr. R.J. van Kemmenade summarizes the novel biomarkers in Heart Failure.87

The Need to look for a Selective MMP Inhibitor

Since Weber et al identified the Renin-AngiotensinAldosterone System (RAAS) with eventual myocardial fibrosis and stiffness in 1994 we have treated most forms of heart failure thus far with Diuretics, Anticoagulants, Digoxin, Beta-Blockers, ACE Inhibitors and Aldosterone Inhibitors.88-90 We have recently discovered that inflammatory markers inducing MMP activation are in play even before the RAAS system is activated in heart failure.91-92 Therefore, MMP inhibition could be considered as a potential early intervention strategy. Positive results depicting a decrease in collagen deposition have been recorded in animal trials yet the transfer to human trials has been slow due to the side-effects brought about by the many beneficial and vital functions of the MMPs.93 Selectively identifying an MMP causing pathological remodeling without a concomitant role in health has proven difficult with limited efficacy in the former and significant musculoskeletal side-effects in the latter.94 Nevertheless, the future of early treatment in patients with heart failure could very well lie in treating their specific matrix pathology, and not in purely treating the neurohormonal systemic symptoms, as is current practice.
### Table 2: The Different Etiological Subtypes of Cardiomyopathy along with Their Mechanism, Biomarkers and Clinical Features based on the specific condition and body.

<table>
<thead>
<tr>
<th>Mechanism</th>
<th>Ischemic</th>
<th>Idiopathic</th>
<th>Hypertrophic</th>
<th>Hypertensive</th>
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<tr>
<td></td>
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<td>Inflammatory Cascade</td>
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<td></td>
<td>D TIMP-1, D TIMP-2,</td>
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<td>Vilahur et al, 2012</td>
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<tr>
<td>Features</td>
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<td>Hypertrophy (matrix and</td>
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<td></td>
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<td>Systolic Failure</td>
<td>Diastolic Failure</td>
<td>Perivascular Collagen</td>
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</tbody>
</table>

**Legend:** MMP – Matrix Metalloproteinase, TIMP – Tissue Inhibitor of Metalloproteinase.

### Personalizing Heart Failure Treatment to each Patient

Miller and Miller suggested that the future of heart failure medicine will be tailoring treatment to each patient’s specific condition and body. As Table 2 describes, there are many different routes to acquiring heart failure and they will all likely produce very different inflammatory responses to disease with differing amounts and type of collagen remodeling. As a result of relatively recent research, we now know that collagen consistency will widely differ amongst the population in the amount of cross-linked collagen fibers they will exhibit, affected by disease and age. In a canine trial, Jagquist showed that age affected an animal’s response to heart failure medication post-MI. Also, as Table 3 will highlight, different causes of heart failure all vary in their exact mechanism and pathophysiology and therefore require specific and individual treatment. Following a one-fits-all treatment plan might not be the most successful approach considering the vast amount of research that has been done on the cardiac ECM until now.

### Conclusion

Heart Failure is a complex and progressive disease involving both the cells and ECM of the myocardium. A tip in the fine balance of ECM dynamic remodeling, mostly concerning MMPs and TIMPs, affects the heart’s physiological properties immensely and can lead to the dysregulation that provides the basis for progression to heart failure in the future. These imbalances within the myocardium occur through distinct mechanisms of variable root cause, as shown in Table 2, with the final result remaining the breakdown of a stable matrix and the common pathophysiology that will ensue. The current research field requires further clarification into the key protein modulators that have an effect on the ECM in disease, other than the enzymatic MMPs and TIMPs. Although many discovered, it is yet unclear as to which changes are primary or secondary to the remodeling process. Zamplpa and Lindsey concluded with a similar view in the need for a catalogue on the specific changes that occur in the ECM with cardiac injury. Many studies have pointed out Osteopontin, Galectin, Periostin and Integrin as potential modulators. The near future will help us get an even clearer picture of the roles and importance of these non-collagenous proteins in heart failure. The field of the Cardiac ECM holds many academic and clinical applications and we have yet to realize its true potential in combatting the chronic issue of heart failure.
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Acknowledgments
I acknowledge the advisory help received from Professor of Cardiac Pathophysiology Andrew Trafford (BVSc, CertIVA, PhD, MRCS). There was no funding or endorsement from any third party. All the work shown is truthful, objective and with no agenda on mine or my Supervisor’s behalf.

Conflict of interest statement & Funding
No conflict of interest exists for this article. No financing was involved for this article.

Author contributions

Cite as:
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