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Antioxidant Anthocyanidins and Calcium Transport Modulation of the Ryanodine Receptor of Skeletal Muscle (RyR1)

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Antioxidant Anthocyanidins and Calcium Transport Modulation of the Ryanodine Receptor of Skeletal Muscle (RyR1)

by

Thomas J. Dornan

A thesis submitted in partial fulfillment of the requirements for the degree of

Master of Science

in

Physics

Thesis Committee:
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Abstract

Cardiovascular disease (CVD) claims more lives than any other disease in the world. Although numerous biological pathways share the blame, ventricular tachyarrhythmia (VT) is estimated to account for ~25% of all CVD deaths. A complete understanding of the molecular mechanisms underlying VT is unknown but recent studies have linked VT to improper calcium handling in the heart (canine). The principle calcium regulator in the muscle cell is the calcium ion release channel (aka RyR). Numerous endogenous and exogenous compounds can affect the way the RyR regulates calcium. In particular, abnormal levels of oxidants (reactive oxygen species) can oxidize critical thiol groups on the RyR and modulate its activity. Interestingly, high levels of oxidants are also associated with numerous bodily disease states including cancers, muscle fatigue/failure, and CVD.

In this thesis, two important dietary antioxidant compounds, the anthocyanidins pelargonidin and delphinidin, are evaluated for their effects on regulating the transport of calcium through the calcium release channel (RyR1) of the sarcoplasmic reticulum of skeletal muscle. Pelargonidin and delphinidin are structurally similar with delphinidin only differing from pelargonidin by the addition of two hydroxyl groups. Both compounds undergo time dependent structural changes in aqueous solutions at physiological pH and a mixture of more than
four structures of each compound can be present in solution simultaneously. Pelargonidin and delphinidin show distinct differences in their calcium flux regulating effect on the RyR1. Delphinidin stimulates calcium flux and RyR1 activity whereas pelargonidin can cause both inhibition and stimulation of the RyR1. The strength of stimulation and inhibition of calcium transport through the RyR by delphinidin and pelargonidin may be attributed to the structural and chemical changes in those compounds that occur in solutions near physiological pH and the subsequent chemical characteristics of the diverse set of structures that are simultaneously present in solution.
Dedication

This thesis is dedicated to my mother and father. Their unconditional love and support throughout the years has shaped my soul and provided me with all of the opportunity that I've ever needed. They both passed away during the course of this thesis work but I can easily see the smiles on their faces in response to this achievement.
Acknowledgements

A sincere appreciation is extended to my advisor Jon Abramson for all of his time and support with regards to the completion of this project. Additionally, financial support was partially provided by the award of a teaching assistantship for two years by the Department of Physics and by the award of a two term research assistantship provided by Erik Sanchez in conjunction with Jon Abramson.
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Cardio vascular disease (CVD) causes more deaths than any other disease on the globe (World Health Organization). Although CVD encompasses numerous distinct pathological mechanisms, sudden cardiac death (SCD), the abrupt loss of heart function, is estimated to account for approximately 50% of CVD related deaths. A majority of the SCD fatalities is attributed to ventricular tachyarrhythmias (VT) (Huikuri, Castellanos, and Myerburg 2001). In an effort to improve the quality of life for non-fatal CVD victims, reduce CVD deaths, and minimize the enormous economic costs associated with CVD, extensive epidemiological studies and laboratory research continue to be funded and performed in hopes to better explain the pathological molecular mechanisms of CVD.
Non-behavioral risk factors of CVD include increasing age, male gender, and heredity related factors (children of people with heart disease and race related increased risks). Behavioral risk factors (ones that we can change) account for over 90% of the risk of heart failures (HF). These factors include cigarette smoking, abnormal blood lipid levels (high cholesterol), hypertension (high blood pressure related), diabetes, abdominal obesity, a lack of physical activity, low daily fruit and vegetable consumption, alcohol over consumption, and psychosocial index. (AHA-Heart and Stroke Update 4/2009 referenced Lancet. 2004;364:937–952).

Interestingly, a subsequent epidemiological study found an exception to that rule, the people of southern France consume more fat and smoke more cigarettes than other countries with a similar incidence of CVD deaths. Additional studies revealed that the same population also had a slightly different diet which included higher intakes of vegetables, fruits, and wine. The lower incidence of CVD was quickly attributed to the constituents of those foods which were abundant in flavonoid antioxidant compounds. Since then, research has been conducted on almost every conceivable natural antioxidant to be found in those foods. Extensive quantitative-structure activity relationship (QSAR) studies have characterized those compounds for their antioxidant properties. Naturally, those compounds that exhibited the highest level of antioxidant activity in vitro were perceived as the best of the best’. These compounds have been studied extensively in the literature. If antioxidants are the ‘cure’, then it follows that
oxidants may be the cause. Researchers targeted native reactive oxidant species (ROS) and studied their effects on the proliferation of CVD biomarkers in vitro. The results all seem to imply that ROS cause severe damage to biological processes and can be attributed to increased incidence of several CVD pathologies. In addition to the ROS connection, recent studies (Lehnart et al. 2004; Belevych et al. 2007; Belevych et al. 2008; Belevych et al. 2009; Belevych and Terentyev 2010) related to ventricular tachyarrhythmias, a significant contributor to sudden heart failure, have shown a connection between VT and increased levels (overload) of calcium (Ca^{2+}) in the heart muscle cell via ‘leaky’ calcium release ion channels (ryanodine receptor). The ryanodine receptor (RyR) is known to have reactive thiol residues that are extremely sensitive to the oxidation by ROS and numerous other compounds. Oxidation of these thiols can causes increased channel activity and Ca^{2+} conductance (J.J. Abramson and G. Salama 1988; Jonathan J. Abramson and Guy Salama 1989; J. J. Abramson et al. 1988; J. J. Abramson, Cronin, and G. Salama 1988).

There may indeed be a connection between increased oxidative stress in the cell (oxidative stress) and abnormal calcium handling by the RyR (calcium overload) in cardiac myocytes, and deaths by ventricular tachyarrhythmias. Dietary antioxidants that play a part in reducing ROS in the cell may indeed have part in the decreased incidence of CVD deaths of those populations engaged in higher (disease) risk activities. However, numerous studies have shown that antioxidant compounds, especially the good ones, can also act as prooxidants. These
antioxidant compounds have also been shown to bind to proteins and protective enzymes impacting their function (Graziani et al. 1982; Kaldas et al. 2005). The intention of the investigation of various dietary antioxidants is to characterize that compound for its potential beneficial properties and its potential deleterious properties so that fully informed and educated decisions can be made as to whether or not their benefits outweigh their risks. Due to the possible prooxidant activity of flavonoid antioxidants, it is clear that before recommendations can be made regarding their dietary inclusion, any impact that they may have on other biological systems must be understood. Given the importance of Ca$^{2+}$ homeostasis in both skeletal and cardiac muscle, the interaction of these compounds need to be evaluated with regards to their possible modulation effects on the RyR.

1.1 Oxidative Stress and Calcium Homeostasis in Muscle

Calcium can signal and activate biological processes via dynamic variations in its concentration within cells. The concentration of free Ca$^{2+}$ is generally controlled by its transport in and out of the cytoplasm of cells by ion channel proteins located in the membrane of the endoplasmic/sarcoplasmic reticulum organelles within the cell. This Ca$^{2+}$ movement alters the cytoplasmic (intra-cellular) Ca$^{2+}$ concentration which in turn impacts Ca$^{2+}$ binding to various biological substrates. An important example of Ca$^{2+}$ signaling is the binding of Ca$^{2+}$ to specific muscle...
cell proteins. This interaction is mandatory for our muscles to contract and it also controls how much force is generated during the contraction. Indeed, without Ca\(^{2+}\) signaling, our muscles would cease to function and we would die. Improper functioning of the Ca\(^{2+}\) regulatory proteins can lead to pathological Ca\(^{2+}\) homeostasis conditions that can severely decrease the quality of human life and also lead to eventual death. The importance of knowing how Ca\(^{2+}\) levels in muscle cells is regulated and knowing what cause changes in that regulation cannot be overstated. Studies to determine why muscles malfunction, fatigue, and fail have consumed researchers for many decades. Although the normal functioning of the muscle is the result of synergistic cooperation from numerous proteins that reside in the muscle cell, the main regulatory protein, the one that controls the fast Ca\(^{2+}\) transient signals within the cell, is called the ryanodine receptor (RyR). The RyR is, simply put, the valve that opens and closes controlling the concentration of Ca\(^{2+}\) within the cell. Increased Ca\(^{2+}\) levels allow for the Ca\(^{2+}\) - protein interactions that allow the contractile process of the muscle to occur. The RyR is very sensitive to the environment within the cell. It is sensitive to Ca\(^{2+}\) concentrations, the redox environment, binding of other closely associated proteins, and numerous other endogenous and exogenous compounds. All of these can have an effect on the way the RyR manages the Ca\(^{2+}\) level in the cell. This is of utmost importance because too much, or too little, Ca\(^{2+}\) can cause the concert of cooperation between the cellular proteins to breakdown which may lead to various pathological conditions and even death. It
has been shown that one of the most important cellular conditions, with regards to proper Ca\(^{2+}\) homeostasis regulation by the RyR, is the redox environment within the cell (Jonathan J. Abramson and Guy Salama 1989; Jonathan J. Abramson and Guy Salama 1988; J. L. Trimm, G. Salama, and J. J. Abramson 1986). The redox potential in the cell is related to the relative ease in which the constituents in the cell environment can reduce (donate electrons or hydrogens to) the RyR or oxidize (accept electrons or hydrogens from) the RyR. The RyR is a large homotetrameric membrane protein containing approximately 5000 amino acids and weighing more than 2.2 megadaltons. The protein contains approximately 320-400 cytoplasmic, oxidation sensitive, cysteine residues (Xu et al. 1998). These cysteines have thiol side chains that are very sensitive to oxidization. When the thiols are oxidized, intra-molecular cystines or cysteine-oxidizer adducts can be formed. It has been shown that some of the thiols on the RyR activate the channel when they are exposed to oxidizing conditions and that some of the thiols inhibit the channel when oxidized. (Jonathan J. Abramson and Guy Salama 1988; Jonathan J. Abramson and Guy Salama 1989; A. Dulhunty et al. 2000). To maintain proper cell function, the redox potential of the muscle cell environment is kept in a tightly controlled and reduced state by the relative ratios (redox couple) of a few important endogenous compounds capable of cyclic redox states. The redox state of the cellular environment can be impacted by all of the cellular redox couples present with each couple contributing via an additive effect. However, due to its high concentration and respective reducing capacity,
the glutathione disulfide/glutathione (GSSG/2GSH) couple is considered to be
the main redox buffer of the cell and the state (value) of this couple alone is
typically used as an indication of the state of the redox environment of the cell
(Schafer and Buettner 2001). The relative concentration of GSH present in the
cell, and consequently, the redox environment is impacted by the invasion of
oxidative species. The main endogenous oxidative species are derivatives of
either oxygen or nitrogen. In particular, oxidative free radical compounds,
hydrogen peroxide, and derivatives of nitric oxide. These are commonly referred
to as reactive oxygen species (ROS) or reactive nitrogen species (RNS). ROS
are natively generated and released from the oxidative phosphorylation
processes of the mitochondria in the cell and by other mechanisms such as
xanthine oxidase (XO) and NAD(P)H oxidases. ROS can impart their damage by
further increasing ROS levels via autoxidation reactions that further increase the
levels of oxidants by directly attacking and altering (taking or donating electrons)
cellular membranes and proteins causing altered function (Sen and Packer 2000).
Increased levels of ROS can be generated during increased levels of exercise,
exposure to environmental pollutants, and consumption of pro-oxidant
compounds in food and beverages. In order to keep the redox environment
tightly controlled (and in a reduced state), the body has a host of endogenous
antioxidant defenses against these oxidants. Antioxidants are tasked with
neutralizing the ROS by donating or accepting electrons or hydrogens to or from
the ROS minimizing or nullifying its reactivity before the ROS reaches critical
cellular substrates. In doing so, the antioxidant generally becomes a radical species itself but one that is more stable and relatively unreactive as compared to the original ROS. The main endogenous antioxidant compounds of the cell are glutathione (GSH), the glutathione peroxidase system, CoEnzyme Q, superoxide dismutase (SOD), and catalase. This defense network is effective at protecting against various ROS under normal cell conditions but high levels of oxidants, generated during times of high oxygen consumption (strenuous exercise) or as a result of prooxidant intake (air pollutants or high fat diets), can overwhelm the endogenous defenses. These increased levels of oxidant compounds can easily overwhelm the endogenous antioxidants inducing a state of oxidative stress in the cell (M. J. Davies and Dean 1997; Fridovich 1978; K.J. Davies et al. 1982).

1.2 Oxidative Stress and Disease

It is now well accepted that elevated ROS levels are associated with numerous bodily disease states including numerous cardio vascular diseases including but not limited to atherosclerosis pathology (Khatri et al. 2004), cardiac hypertrophy (Higuchi et al. 2002), and myocardial lipotoxicity (S. Sharma et al. 2004). It is also connected with a variety of cancers, Alzheimer’s Disease, inflammatory diseases (Hidalgo and Donoso 2008; Zima and Blatter 2006; Belevych et al. 2009) as well as skeletal muscle fatigue and failure (Moopanar and David G Allen 2005). Lipid peroxidation and sulfhydryl group oxidation during ischemia-
reperfusion can lead to abnormally high cell Ca\(^{2+}\) levels and contractile
dysfunction (Dhalla et al. 1999). Improper Ca\(^{2+}\) handling in the heart muscle has
been connected to atrial arrhythmogenesis and contractile dysfunction in dogs
with congestive heart failure (CHF) (Yeh et al. 2008). Evidence has shown that
reactive oxygen species (ROS) alter calcium pump function (Kourie 1998; Kaplan
et al. 2003), alter calcium current reduction in L-type calcium channels (Guerra et
al. 1996), and decrease contractility via altered calcium handling (He et al. 2003)
in cardiac myocytes. Recently, there have been numerous studies pointing to
Ca\(^{2+}\) overload via improper Ca\(^{2+}\) handling by the RyR as an underlying cause of
arrhythmias and sudden cardiac death (Belevych et al. 2007; Belevych et al.
2009).

1.3 The Flavonoid Antioxidant – Calcium Homeostasis Connection

![Figure 1.2 Structure of Quercetin](image)
The effects of flavonoid compounds on the activity of the RyR need to be evaluated before they can be considered viable nutraceuticals. These compounds may be good antioxidants and even promote anticancer effects but if they or their radical species react directly with the RyR or other Ca$^{2+}$ homeostasis associated proteins, they may cause harmful calcium overloading effects.

Studies on the interaction of flavonoids with the RyR are sparse. However, one popular flavonoid antioxidant, quercetin, has been shown to activate Ca$^{2+}$ release from the SR skeletal muscle (D. H. Kim, Ohnishi, and Ikemoto 1983). Quercetin is a popular dietary antioxidant found in numerous vegetables, fruits, nuts, and teas. It has been extensively studied due to its excellent antioxidant properties and inclusion in numerous foods. Lee et. al. (Eun Hui Lee, Gerhard Meissner, and Do Han Kim 2002) showed that quercetin activates single RyR channel open probability ($EC_{50}$ of 91.0 +/- 15 µM at 10 µM Ca$^{2+}$) more effectively than the well known RyR activator caffeine ($EC_{50}$ of ~2.93 mM at 10 µM Ca$^{2+}$). Quercetin also exhibited a higher maximal open probability ($P_o$=0.89 at 250 µM) as compared to caffeine ($P_o$=0.66 at 9 mM). Quercetin is effective at stimulating the RyR but it didn’t exhibit any inhibitory actions. With regards to the redox sensitivity of the RyR, further tests were conducted to assay the effects of quercetin in the presence of the reduced and oxidized forms of glutathione (a cellular redox couple and as well as an endogenous antioxidant). The results showed that the effects of 5 mM GSSG (a RyR1 stimulant) plus quercetin on the RyR1 were not reversed by mM GSH (a known RyR1 inhibitor (Anthony C Zable, Terence G
Favero, and Jonathan J Abramson 1997)). This may indicate that quercetin does not work via a redox dependent pathway. They also showed that the known stimulatory effects of K\(^+\)-ATP on the RyR partially inhibited the stimulation by quercetin indicating that they may share the same binding site or that the effects of ATP on the channel may allosterically affect quercetin binding. These results are interesting from the stand point that quercetin appears to have a direct effect on the RyR yet not via a redox pathway. This is surprising since quercetin has a catechol group (ortho hydroxyls) on its B-ring which is notorious for being oxidized to semiquinone radicals and quinones which have prooxidant properties and directly react with protein thiols (Metodiewa et al. 1999; Boots et al. 2003).
1.4 The Anthocyanidins

Anthocyanidins are a subclass of flavonoids and are best known for the beautiful colors they impart to nature. Based on their simplicity and wide ranging abundance, the anthocyanidins delphinidin (Dp), cyanidin (Cy), and pelargonidin (Pg) are considered the three main anthocyanidins of more than 500 that are known to exist. They are responsible for most of the intense oranges, reds, blues, and purple colors exhibited in leaves, flowers, and fruits (J. B. Harborne, T. J. Mabry, and H. Mabry 1975). The chemical and biological properties of these compounds are very complex. Unlike other flavonoids, anthocyanidins are structurally unstable in weakly acidic to alkaline solutions. Upon exposure to these conditions, they will immediately start to transform into several different structural forms each with distinct chemical properties. The relative proportions of these forms is dependent on the pH of the solution and the specific...
functionalization of the compound (Brouillard and Delaporte 1977; Brouillard and Dubois 1977; Brouillard, Iacobucci, and Sweeny 1982; Jurd 1963; Jurd and Geissman 1963; Jurd 1964; Jurd and Waiss 1965) (for more information see chapter 3). Along with the possible health benefits previously mentioned, these anthocyanidins are interesting and commercially valuable due to their ability to act as food colorants and food preservatives (antioxidants).

1.4.1 Pelargonidin and Delphinidin – The Antioxidants

![Pelargonidin and Delphinidin Structures](image)

Figure 1.4 Structures of pelargonidin and delphinidin
Although both pelargonidin (Pg) and delphinidin (Dp) exhibit good antioxidant properties as compared to the well accepted antioxidant vitamin E, delphinidin is dominantly superior to pelargonidin as an antioxidant. Delphinidin’s antioxidant quality is attributed to its tri-hydroxyl structure in its B-ring. Those groups are very easily oxidized and its mechanism of antioxidant action is probably by electron donation and formation of a semiquinone moiety. The donation of a second electron then will form a quinone. (S. A. Van Acker et al. 1996; A. Sekher Pannala et al. 2001). Pelargonidin only has a mono-hydroxyl B-ring and is a less effective antioxidant. Its mechanism of action is probably by donation of an electron and subsequent formation of a phenoxyl radical (Galati et al. 1999; Ananth Sekher Pannala et al. 2001).

Both Pg and Dp exhibit superoxide and hydroxyl scavenging activity and can inhibit hydrogen peroxide induced lipid peroxidation. In a study of pomegranate extract, a hypoxanthine-xanthine oxidase superoxide generating system where DMPO-OOH spin adducts (indicator of superoxide concentration) were measured by ESR, Dp and Pg both exhibited scavenging properties with Dp being over two orders of magnitude better than Pg (IC$_{50}$=0.8µM and 130µM respectively). They also scavenged hydroxyl radicals with Dp being more effective than Pg although their scavenging properties may involve indirect rather than direct scavenging and possibly by more than one scavenging mechanism (Noda et al. 2002).
Dp and Pg also were shown to inhibit H$_2$O$_2$ induced lipid peroxidation in rat brain homogenate when H$_2$O$_2$ and Dp and Pg were incubated together with the brain substrate for 60 minutes (IC$_{50}$=0.7µM and 85µM respectively) (Noda et al. 2002). In a similar extract study (grape seed, etc), Dp and Pg also exhibited (with one exception) scavenging properties for superoxide, hypochlorite ion, hydroxyl radical, and linolenic acid with Dp, again, being a better scavenger in all cases. Pg did not exhibit superoxide scavenging properties in this study (Wada et al. 2007).

Another study assessed the ability of Pg and Dp to scavenge the DPPH (1,1-Diphenyl-2-Picrylhydrazyl) radical (4 min incubation in MeOH), prevent MeLo (Methyl Linoleate) emulsion (10% oil in water) hydroperoxide formation after 72 hours (pH~5.5), inhibit Cu initiated hexanal formation from oxidation of LDL (2 hr incubation @ pH7.4), and prevent bulk MeLo (100%) hydroperoxide formation after 72 hours. Generally, Pg and Dp both exhibited antioxidant activity with Dp being a better antioxidant. However, at a low concentration (2.5µM), Dp exhibited prooxidant activity but Pg was still an antioxidant in the LDL assay. They attributed the better antioxidant property of Pg to the fact that it has low polarity and thus may have better access to the lipid peroxidation site of the LDL (they sited similar results from another study). The researchers also admitted that the method of antioxidant action at higher concentrations may be due to hydrogen donation to the tocopheryl radical back to tocopherol (vitamin E) or by scavenging radicals and protecting tocopherol from consumption. Additionally,
the anthocyanidins may bind with apolipoprotein B and prevent its subsequent binding to LDL thus blocking Cu catalyzed oxidation. Interestingly, they also noted that polar antioxidants (hydrophilic) are generally more efficient in bulk oils than non-polar antioxidants (hydrophobic). This is due to a higher concentration or accumulation of a hydrophilic compound that accumulates at the oil-water interface (Kahkonen and Heinonen 2003).

The radical scavenging ability of Pg and Dp is pH dependent. In a study by Borkowski et al., the TEAC antioxidant values (antioxidant action relative to Trolox) are measured for Pg and Dp (and several other anthocyanidins). The TEAC assay is based on the ABTS$^{+}$ radical and the ability of the antioxidant to scavenge it and thus reduce its characteristic absorption spectrum. This particular compound was used due to its stability in a wide range of pHs (2-9.5). Results basically indicated that Dp is a powerful antioxidant (several times better than Trolox) and its increase in scavenging activity is related to the deprotonations of its C4’ hydroxyl ($pK_a$ ~5.3 (experimentally determined)) and its C5 hydroxyl ($pK_a$ ~6.73 (computationally predicted)). Pg on the other hand had an activity similar to Trolox but did not show a strong increase in activity with the deprotonation of C5 ($pK_a$ ~5.79 (experimentally determined)) and the deprotonation at C4’ ($pK_a$ ~7.05 (computationally predicted)) (Borkowski et al. 2005).
1.4.2 Pelargonidin and Delphinidin – The Prooxidants

As hinted at in the previous section, both Pg and Dp can exhibit prooxidant properties. Delphinidin’s pyrogallo group in its B-ring is highly oxidizable and autoxidation of this group is possible at physiological pH. Autoxidation is accompanied by generation of superoxide and hydrogen peroxide (Hodnick et al. 1986; Hodnick et al. 1988). This property may also be enhanced by delphinidin’s metal chelating ability which is also known to catalyze the autoxidation of (similarly oxidizable) catechol groups (dihydroxy group) (Vanacker et al. 1995; Canada et al. 1990; Dangles, Fargeix, and Dufour 2000).

Pelargonidin, with only a monohydroxyl group in the B-ring, does not exhibit autoxidation and doesn’t chelate metals. Its reduction in hydrogen peroxide induced lipid peroxidation activity may be due to a different oxidation mechanism (Saskia A. B. E Van Acker et al. 1996). In one study, it was found that the phenoxy radicals of similarly structured compounds with a mono-hydroxyl substituted B ring redox cycled and caused extensive GSH oxidation at pH~7 to form thiyl- radicals which reduced oxygen. Surprisingly, flavonoids containing a catechol B-ring did not show the same oxidative properties (Galati et al. 1999).

In a LDL oxidation assay (Satue-Gracia, Heinonen, and Frankel 1997), both Dp and Pg act as prooxidants when 3uM copper is used as an oxidation initiator. Interestingly, when 10µM copper is used, Pg acts as an antioxidant and Dp continues to be prooxidant. It was noted that more work was needed to be able
to propose a mechanism for why Pg changes its antioxidant/prooxidant activity with various copper concentrations.
1.5 This Thesis

Due to the importance of understanding the effects that potential nutraceuticals have on the Ca$^{2+}$ homeostasis in muscle tissue, this thesis will attempt to evaluate the modulation effects that pelargonidin and delphinidin have on the ryanodine receptor calcium release channel of skeletal muscle.
2 Muscle Structure and Function

At a macro level, muscles are well defined organs capable of generating great force when stimulated to do so. Some of these stimuli are consciously controlled (voluntary muscle) and some are not consciously controlled (involuntary muscle). The heart (cardiac muscle), the iris of the eye, the arrector pili of the skin, the bladder, blood vessels, the gastrointestinal tract, are all examples of organs that contain involuntary muscle. Skeletal muscle is the only muscle tissue that is consciously controlled. It is responsible for locomotion, posture, voluntary eye lid movement, and sphincter control to name a few. Both cardiac muscle and skeletal muscle are classified as striated muscle due to the striated appearance of muscle fibers when viewed under the light microscope. These striations are due to a repeating, end-to-end, pattern of contractile proteins at the molecular level of the tissue. There are three types of muscle tissue: cardiac, skeletal, and smooth. They all have unique characteristics depending on the function for which they are designed. For instance, non-striated smooth muscle lacks the end-to-end pattern of contractile proteins as mentioned. The random orientation and alignment of the contractile proteins of smooth muscle allows it to accommodate larger variations in stretch while still maintaining contractility. These characteristics are favorable in organs such as the bladder and the GI tract. On the other hand, the end-to-end and aligned orientation characteristics of skeletal and cardiac muscle allow them to generate large forces with longer contractile
distances. These characteristics are required for generating large movements like bending a leg and also for generating high fluid pressures within the heart ventricles.

2.1 Muscle Structure

The original work presented in this thesis is most closely related to the function of the calcium release channel of skeletal muscle and thus it will be the primary muscle tissue discussed. However, probably the most benefit to society can be obtained if these types of studies are applied to cardiac muscle. For that reason, the finer details of the function of cardiac muscle may also be discussed. Smooth muscle is also very important with regards to human anatomy and physiology, but due to the limited scope of this thesis, it will generally not be discussed.

2.1.1 Skeletal Muscle Macro Structure

Skeletal muscle is composed of both muscle tissue and connective tissue and is highly compartmentalized. Each compartment is contained and protected by fibrous connective tissue (epimysium, perimysium, and endomysium). The largest of these compartments contains the whole macro muscle (muscle belly) and is surrounded by the epimysium. The muscle belly is composed of fascicles which are many smaller bundles of parallel strands encased in the perimysium. The fascicles contain groups of the long muscle cell which also known as the myofiber.
2.1.2 The Muscle Cell Structure

Skeletal myofibers span the complete length of the bulk muscle that contains it. They range from 10 µm to 100 µm in diameter and up to 30 cm long (depending on the macro muscle length). Due to their long shape, myocytes are also referred to as myofibers. Myofibers are surrounded by the endomysium which provides the fiber access to capillaries and nerves. Layered underneath and proximal to the endomysium is the cell membrane of the myofiber, the sarcolemma. The sarcolemma is the plasma membrane that contains the intracellular fluid of the myofiber called the sarcoplasm. The sarcolemma is designed to receive nervous system stimuli and conduct that signal throughout the cell via membrane invaginations (transverse tubules or T-tubules) that project directly into the cylindrically shaped cell at locations associated to the Z-disks of the sarcomere (contractile sub-unit of the myofiber). Internal to the sarcolemma are highly aligned and orientated filamentous motor proteins (myofibrils). Surrounding the myofibrils throughout the myofiber resides an organized and highly functional organelle network called the sarcoplasmic reticulum (SR). The SR is designed to act as a storage compartment for Ca\(^{2+}\). The SR extends from T-tubule to T-tubule and the group of organelles where two SR membranes from adjacent sarcomeres meet a t-tubule is called a triad (SR membrane: T-tubule: SR membrane). Proximal to the T-tubule, the SR exhibits an enlargement or sac which is referred to as the terminal cisternae (TC). The matrix of SR between the T-tubules and proximal to the longitudinal center of the myofibrils is the
longitudinal SR. The myofiber is composed of an end-to-end chain of many contractile sub-units called the sarcomere. The sarcomere is a highly organized and complex arrangement of numerous proteins. Filamentous actin (f-actin or thin filament) and filamentous myosin (f-myosin or thick filament) are the contractile proteins of muscle. Tropomyosin and troponin are the primary regulatory proteins. Additionally there are structural proteins that help maintain the form of the muscle cells. Titin is an elastic protein filament that runs through the center of the myosin filaments and emerges from the end of the filament to connect to the Z-disks proximal to each end of the filament. The Z-disks are composed of an elastic protein filament called connectin. The Z-disks are the highly light refracting portion of the fiber that gives rise to the striated appearance of the muscle. Z-disks provide direct anchorage for both the titin and the actin filaments. The area between two Z-disks that contains both f-actin and f-myosin is the sarcomere.

2.1.3 The Sarcomere

F-myosin filaments (thick filaments) are about 15 nm in diameter and are made of hundreds of motor proteins molecules, myosin II (hence forth referred to as just myosin). One molecule of myosin is comprised of two polypeptide chains intertwined to form a golf club shaped structure where one end is the tail and the other end the head. A myosin filament is several hundred of these golf club shaped molecules linked together end to end with the tail sections intertwined and the heads extending outward from the filament. Thin filaments are about 7
nm in diameter and are made up of two twisted strands of f-actin (filamentous actin). F-actin is a string like assembly of globular actin (G-actin) protein molecules. Each G-actin molecule has an active site that can bind to the head domain of the myosin molecules of F-myosin. Wrapped around the thin filament is another long polypeptide chain of proteins called tropomyosin. Tropomyosin has the function of blocking the binding of the myosin heads to the G-actin. Bound to the tropomyosin is a smaller and intermittently spaced calcium binding protein called troponin. Troponin has the function of binding calcium, undergoing a conformational change, and moving the tropomyosin filament which allows myosin to bind to the G-actin sites. Troponin has three sub-units and is bound to both the thin and thick filaments. The three sub-units are: TnI (bound to actin), TnT (bound to myosin), and TnC (calcium binding sub-unit).

When the sarcomere is viewed under a light microscope, the varying densities and refracting properties of the protein structures can be observed, and thus distinguished. The area spanning adjacent sarcomeres that includes the Z-disk but does not include the F-actin is called the “I Band”. The area that spans the length of one F-myosin filament is called the “A Band”, the area near the center of the F-myosin that does not overlap F-actin is called the “H zone”, and the area where the F-actin meets at adjacent sarcomeres is connected is called the Z-disk (as explained previously). The length of a sarcomere is approximately 1.7 µm in the contracted state and 2.2 µm in the extended state. The F-myosin’s structure is such that the myosin heads orient themselves in a corkscrew shape along the
filament. This gives rise to a repeating pattern with six crossbridge head groups positioned in a radial pattern at multiples of 60 degrees. These repeating patterns are oriented such that the crossbridge heads align with each other along the length of the F-myosin. F-actin and F-myosin are orientated in hexagonal pattern when viewed as a cross section. There are six F-actin filaments surrounding each F-myosin filament, one F-actin per crossbridge head row and one F-actin filament may bind to more than one F-actin filament. This orientation is maintained throughout the myofiber which contains hundreds of myofilaments per fiber.

2.1.4 The Sarcoplasmic Reticulum (SR)

The SR is an organelle in the myocyte that has the primary function of storing $\text{Ca}^{2+}$. The SR is a webbed network of an enclosed membranous system extending along the length of each sarcomere. As previously noted, the SR has two main structural and functional regions: 1) the terminal cisternae are the enlarged sac like structures nearest to the T-tubules and 2) the longitudinal SR is located along the length of the sarcomere. The area where the TC meets the t-tubule is highly functional and complex and it's commonly referred to as the triadic junction, the dyadic junction, or (when referring to the SR specifically in this region) the junctional SR. The TC region of the SR contains a large amount of the $\text{Ca}^{2+}$ binding protein calsequestrin. Calsequestrin is able to bind up to 50 $\text{Ca}^{2+}$ ions per monomer and this binding of free $\text{Ca}^{2+}$ increases the total $\text{Ca}^{2+}$ storage potential of the SR. The junctional SR is the home of the main regulatory
Ca$^{2+}$ ion channel, the ryanodine receptor (RyR) (see next section more for additional detail) as well as other associated transmembrane proteins such as junctin and triadin. On the T-tubule, there resides a voltage gated Ca$^{2+}$ ion channel called the dihydropyridine receptor (DHPR) which is also an important player in the functionally of muscle. Located primarily in the longitudinal SR is the Ca$^{2+}$ ATPase pump (aka SERCA). There are almost 100 fold more SERCA pumps than RyRs in the SR. The Ca$^{2+}$ ATPase is responsible for accumulating Ca$^{2+}$ into the SR which allows the muscle to relax.

2.1.5 Muscle Contraction

The signal for a muscle to contract is sent from the brain via a neuron/axon to the neuromuscular junction. There, the signaling action potential opens voltage gated Ca$^{2+}$ ion channels in the axon’s plasma membrane which causes the internal levels of Ca$^{2+}$ to increase. The elevated Ca$^{2+}$ causes intracellular vesicles containing acetylcholine (ACh) to fuse with the cell membrane and, through exocytosis, dump the Ach into the synaptic cleft. ACh then binds to ligand gated Na$^+$ ion channels causing them to open and allow the subsequent influx of Na$^+$ into the cell. This process causes the immediate opening of K$^+$ ion channels and the subsequent efflux of K$^+$ out of the muscle cell. Because of concentration differences, more Na$^+$ flows into the cell than K$^+$ flows out of the cell and this causes an electrochemical depolarization across sarcolemma. This
depolarization self-propagates over the whole muscle cell and down the T-tubules. The depolarization of the T-tubules signals for the opening of the resident voltage gated L-type Ca\(^{2+}\) ion channel, the dihydropyridine receptor (DHPR). In the skeletal muscle cell, the DHPR is believed to be mechanically coupled to the ryanodine receptor and the opening of the DHPR has shown to be the determinate cause of the opening of the ryanodine receptor. In cardiac muscle, the DHPR is not structurally associated with the ryanodine receptor. The opening of the DHPR allows Ca\(^{2+}\) to enter the cell from the cytoplasm. The process of Ca\(^{2+}\) influx through the DHPR and subsequent elevation in Ca\(^{2+}\) concentration is responsible for cardiac ryanodine receptor activation. This process is called “calcium induced calcium release”. The elevated levels of cytoplasmic Ca\(^{2+}\) causes Ca\(^{2+}\) binding to troponin which moves tropomyosin and allows myosin access to the actin binding sites. Myosin binding to the G-actin initiates the conformation change of myosin from a high energy conformation to a low energy conformation. This is commonly referred to as the power stroke. This change pulls the thin filament along the thick filament. The thin filament is attached to the Z-disks on each end of the sarcomere and the power stroke causes the sarcomere to shorten or contract. Thousands of these cellular contractions occurring in unison causes the muscle contraction we’re familiar with at the macro level. After the power stroke occurs, the low energy conformation of myosin remains attached to f-actin until an ATP molecule binds to it. If there isn’t any ATP available, the muscle remains in this contracted state.
(rigor). If ATP is available, it binds to the myosin head and the head de-binds from actin. Once it is free, myosin can hydrolyze the ATP into ADP and $P_\text{i}$. The hydrolysis of ATP provides the energy for myosin to change back into the high energy conformation and then rebind to actin. This cyclic process is referred to as cross-bridge cycling. When the signal from the central nervous system stops initiating depolarizations of the cell membrane, the DHPR closes and the RyR closes as a result. $\text{Ca}^{2+}$ concentrations are decreased and troponin changes its conformation to allow tropomyosin to block actin-myosin binding. This is the resting state of the muscle.

2.1.6 Excitation-Contraction Coupling

Excitation-Contraction Coupling (EC coupling) is the term used when describing the process of converting the cell depolarization into the actual muscle contraction. Although the overall process of the sliding filaments (sliding filament theory) is well established and accepted, the mechanism in which the DHPR communicates the depolarization signal to the RyR in skeletal dyads still remains uncertain. Cardiac dyadic junctions are organized differently than skeletal dyadic junctions. RyRs have the ability to self-organize into repeating patterns and do so in the junctional SR of both cardiac and skeletal muscle. Cryo-EM images of dyads have shown that the DHPRs of cardiac dyads are not arranged in any pattern with respect to the RyRs. They oppose the RyRs but in a random way. However, the DHPRs of skeletal dyads are arranged so that every other RyR1 is directly opposed to a cluster of four DHPRs and each cluster is associated with
one monomer of the RyR (Block 1988). It has been shown that cardiac RyRs don’t require the interaction with the DHPR to release large stores of Ca\textsuperscript{2+} from the SR. Instead, fast increases of Ca\textsuperscript{2+} near the junctional SR causes Ca\textsuperscript{2+} release from the RyR. For this reason, the release of Ca\textsuperscript{2+} from the SR of cardiac SR has been characterized as calcium induced calcium release (CICR). For skeletal dyads, the mechanism is more complex. The highly organized arrangement between the DHPR and the RyR implies a high level of association between them and indeed it has been shown that in functional triads, the molecular modification of what is called the II-III loop or the DHPR can cause the RyR to cease to release Ca\textsuperscript{2+} from the SR. Studies of simpler protein complexes like the voltage dependent K\textsuperscript{+} ion channels have shown that cell membrane depolarizations trigger movements in voltage sensing domains and those movements can be trans-located across the membrane (Jiang, A. Lee, et al. 2003; Jiang, Ruta, et al. 2003). Together with the organized DHPR-RyR1 structures, it is hypothesized that charge dependent movements in the DHPR directly interact with the RyR and trigger the opening of the channel causing Ca\textsuperscript{2+} release. However, there are still a lot of unanswered questions stemming from a wealth of research and discussion and the understanding of the exact mechanisms required for EC coupling is still undetermined.

### 2.2 The Ryanodine Receptor

Ryanodine receptors are the largest membrane ion channels known. Three mammalian RyR isoforms are known to exist. These include the skeletal muscle
RyR (RyR1), the cardiac muscle RyR (RyR2), and the smooth muscle RyR (RyR3). All of the isoforms have a sequence homology of approximately 70%. The skeletal RyR1 (rabbit) sequence consists of 5,037 amino acids and has a molecular weight of 565,223 Da. The complex is a homotetramer of four identical subunits with a total weight of ~2.3 MDa.

### 2.2.1 The Ryanodine Receptor – Structure

Knowing the 'exact' structure of a dynamically functional protein can give vast insight as to the biophysical mechanics and chemistry of the molecule. Unfortunately the RyR is too small to be viewed under a light microscope and too large and complex to solve with traditional structural techniques such as x-ray crystallography. Original freeze-fractured electron microscopy images of skeletal triads by Ferguson et al. (Ferguson, Schwartz, and C Franzini-Armstrong 1984) were among the first to show the square foot structure of the skeletal RyR. They also were able to show the significant association of the RyR and DHPRs in skeletal muscle (Clara Franzini-Armstrong and Kish 1995). Since then, cryo-EM of single particles has emerged as a very useful technique to study the structure of these particles. Utilizing fluorescent probes with EM can also give us an idea of where important modulators bind on the macro structure. Sample preparation includes solubilizing the protein with detergent to separate it from the SR and then freezing it in liquid ethane (~ -88°C). Many images (1,000 to 100,000) are computationally aligned and averaged and screened for artifacts to obtain the final result. Since the proteins have many different orientations, images from
different perspectives are obtained and images can be reconstructed to produce 3-D images (Serysheva, Chiu, and Ludtke 2007). Capturing images of different conformational states is also possible by putting the RyR in an environment that favors a particular conformation prior to freezing it. Sub-nanometer resolutions have now been achieved by several groups (Ludtke et al. 2005; Serysheva et al. 2008; Serysheva et al. 2007) and it has been shown by some of these same groups that resolutions of 6-10 Å is adequate for determining secondary molecular structures (α-helices and β-sheets).

2.2.2 Sub-nanometer cryo-EM Structures

The RyR has a mushroom shape with a large cytoplasmic (CY) head and a smaller transmembrane (TM) stem. From above it has four-fold symmetry (quatrefoil shaped). The CY region is 280 Å x 280 Å x 120 Å with the 120 Å being the dimension from the SR membrane to the top of the channel. The CY region is of low amino acid density and shows numerous possible locations for ligand binding. The TM region is much more dense and has dimensions of 120 Å x 120 Å x 60 Å. The orientation of the TM region is rotated approximately 40 degrees with respect to the CY region (Ludtke et al. 2005). At the corners of the CY structure, there are clamp shaped regions. These ‘clamps’ are connected to central rods by dense bridging structures. Images have been captured with the channel in both open and closed states. When going from the closed to the open channel state, conformational changes are apparent with: the removal of amino acid density in the center of the channel (pore opening), CY clamps change from
a closed shape to an open clamp shape, the TM domain rotates relative to the CY domain by ~4 degrees, and mass movements are seen in the central cavity and TM region (Orlova et al. 1996; Serysheva et al. 1999). The binding sites of several channel modulators have been mapped to the RyR structure by fluorescent tagging techniques. Calmodulin (Samsó and Wagenknecht 2002), FKBP12 (Samsó, Shen, and P. D Allen 2006), and other ligand binding sites have been located in the distal CY structure between clamp regions (‘handle’ region). Additionally, RyR mutations associated with malignant hypothermia and central core disease are known to be within the clamp region. This information supports beliefs that the channel is gated by allosteric means.
Figure 2.1 Cryo-EM reconstruction of RyR1 in open and closed states (Orlova, E. et al. Nature Structural Biology v3(6) 547-52. 1996).
2.2.3 The Ryanodine Receptor Function

2.2.3.1 Associated Proteins and Endogenous Modulators of the RyR

Dihydropyridine Receptor (DHPR): The DHPR is an L-type voltage gated Ca\textsuperscript{2+} ion channel located in the T-tubule membrane of the dyadic junction. It is highly associated with RyR1 both from a structural organization relationship and in function. It is the primary Ca\textsuperscript{2+} release initiator and is required for proper RyR1 function. The DHPR is less associated with RyR2. The structural organization is lacking and a direct interaction with the DHPR is not required for RyR2.

FKBP12/12.6: FKBP12 is a 12kDa protein associated with the RyR1 (FKBP12.6 is associated to RyR2) and binds in the ‘handle’ region at a ratio of four FKBP molecules per RyR (one FKBP per monomer). FKBP12 stabilizes the RyR and disassociation of FKBP from the RyR causes an increase in Ca\textsuperscript{2+} conductance.

Calcium: The RyR is a Ca\textsuperscript{2+} activated Ca\textsuperscript{2+} channel. The response of the RyR to free Ca\textsuperscript{2+} concentration is bell shaped. Ca\textsuperscript{2+} conduction through the RyR is minimal at low (<1 µM ) concentrations, maximal \( \sim \) 100 µM (depending on conditions), and again minimal above 1000 µM free Ca\textsuperscript{2+} (J.S. Smith, Coronado, and G. Meissner 1986). This type of behavior is typically attributed to more than one binding site with each binding site having either differing ligand affinities or conformational accessibility limitations. Due to the small size of Ca\textsuperscript{2+}, the accessibility option is probably not a concern. Therefore, the belief is that there is
a high affinity activating binding site that gets populated at low Ca^{2+} concentrations and a low affinity inhibition binding site is populated at high Ca^{2+} concentrations.

**Calmodulin (CaM):** CaM is a Ca^{2+} binding protein with a molecular weight of 16.8 kDa. CaM’s structure is made up of four Ca^{2+} binding domains (EF-hands) and it can bind four Ca^{2+} ions per protein. It is known to bind with RyR1 and cause stimulation at low Ca^{2+} concentrations and inhibition at high Ca^{2+} concentrations. Binding with RyR2 causes only channel inhibition (J. S Smith, Rousseau, and G. Meissner 1989; Tripathy et al. 1995).

**Magnesium (Mg^{2+}):** Mg^{2+} is a known inhibitor of the RyR. Due to its similar nature and size as compared to Ca^{2+}, it is believed that Mg^{2+} may compete with Ca^{2+} for the high affinity inhibition sites which is believed to cause inhibition of the channel with high concentrations of Ca^{2+} (J.S. Smith, Coronado, and G. Meissner 1986).

**Adenosine-5'-triphosphate (ATP):** ATP as well as other adenine nucleotides (cAMP, ADP, AMP, etc) are stimulators of the RyR (G. Meissner, Darling, and Eveleth 1986).

**Glutathione (GSH):** GSH is a tri-peptide of glutamate, cysteine, and glycine. GSH is a good reducing agent and is one of the principle endogenous compounds tasked with keeping the cellular redox status in check. Its concentration in the cell varies between 1 mM and 10 mM. When in an oxidizing
environment, it can be oxidized and form glutathione disulfide (GSSG). GSH (reduced form) inhibits the RyR and GSSG (oxidized form) stimulates the RyR. The two forms show to be competitive and therefore it is believed that they both act on the same regulatory sulfhydryls on the RyR. (A C Zable, T G Favero, and J J Abramson 1997)

**Superoxide (O$_{2}^{-}$):** Superoxide is commonly generated by leaks in the ETC (electron transport chain) pathway of the mitochondria in cells and is a major contributor to oxidative stress. Superoxide has shown to decrease calmodulin content and increase Ca$^{2+}$ efflux from heavy SR. It was also shown that single channel activation was dependent on the presence of calmodulin. (Kawakami and Okabe 1998)

**Hydrogenperoxide (H$_{2}$O$_{2}$):** H$_{2}$O$_{2}$ was shown to induce release from actively loaded SR vesicles at >1 mM concentrations and induce a bi-phasic response in [³H]-ryanodine binding assays (activation range: 100 µM – 10 mM). (T G Favero, A C Zable, and J J Abramson 1995)

**Junctin and Triadin:** Junctin is a ~27kDa MW membrane protein and Triadin is a 95kDa MW membrane protein (Fan, Brandt, and Caswell 1995). Junctin and Triadin both bind to the RyR and are believed to help associate calsequestrin to the RyR (J. M Lee et al. 2004; Goonasekera et al. 2007; Jones et al. 1995; Angela Dulhunty, Lan Wei, and Nicole Beard 2009). They both have been shown
to activate the RyR when added to the luminal side of the channel and seem to activate at independent sites (L. Wei, Gallant, et al. 2009).

**Calsequestrin (CSQ):** CSQ is a 44 kDa Ca\(^{2+}\) binding protein located in the lumen of the SR that effectively binds up to 50 Ca\(^{2+}\) ions per molecule (MacLennan and Wong 1971). CSQ has been found to increase channel activity when removed from the of RyR complex and inhibit activity when added back in to the luminal space (N. A Beard et al. 2002). There are two CSQ genes, CSQ1 (fast and slow twitch skeletal muscle) and CSQ2 (slow twitch skeletal and cardiac). Single channel studies have shown that CSQ1 may inhibit RyR1 but stimulate RyR2 (L. Wei, Hanna, et al. 2009) It should be noted that numerous studies of the interaction between RyR, CSQ, junctin, and triadin are being reported in current literature and important and relevant information regarding the interaction of these protein complexes are being discovered.

**2.2.3.2 Non-Endogenous Modulators**

**Ryanodine (Ry):** Ryanodine is an alkaloid found in the plant Ryania speciosa of South America. It has a high affinity to the RyR (and thus the reason for its name) and locks it in a varying degree open state depending on concentration. (I. N. Pessah et al. 1986; Lai et al. 1988).

**Ruthenium Red:** Ruthenium is a potent channel inhibitor and can block channel activity at sub micromolar concentrations. It is commonly used to verify the presence of the RyR in assays (Ma 1993).
**4-chloro-m-cresol (4-CmC):** 4-CmC a drug preservative stimulates (~75-100 µM) [$^3$H]-ryanodine binding and planar bilayer activity. (Herrmann-Frank et al. 1996)

**Anthraquinones:** Doxorubicin, mitoxantrone, daunorubicin, and rubidaxone all stimulate RyR1 activity by a direct interaction with the caffeine and the Ca$^{2+}$ binding sites. (J. J. Abramson et al. 1988)

### 2.2.3.3 Redox Activity Characteristics

The RyR has approximately 100 cysteines per monomer (Takeshima et al. 1989) with approximately 80 of those accessible to be oxidized. Cysteines are highly reactive and well known to be redox active compounds in other biological molecules. The RyR has also been shown to be very sensitive to its redox environment. When the RyR is subject to an oxidizing environment, the RyR is stimulated, when the oxidizer is pre-treated with a reducing compound (DTT), the oxidizer loses its stimulating capability. (Jonathan J. Abramson and Guy Salama 1989; Jonathan J. Abramson and Guy Salama 1988; Carnes et al. 2009; Marinov BS and J.J. Abramson 2005; Xia, Stangler, and J. J Abramson 2000; Anthony C Zable, Terence G Favero, and Jonathan J Abramson 1997; J.J. Abramson et al. 1983; A. Dulhunty et al. 2000).

Redox active compounds that have shown to modulate the RyR include:
**Heavy Metals** ($\text{Hg}^{2+}$, $\text{Ag}^+$, $\text{Cu}^{2+}$, $\text{Cd}^{2+}$, $\text{Zn}^{2+}$): Heavy metals bind with high affinity to protein sulfhydryl groups and activate the ryanodine receptor at low micromolar concentrations (J. J. Abramson et al. 1983).

**Phtalocyanine Dyes:** Oxidation by the phthalocyanine dyes (alcian blue (AB), $\text{Cu}^{2+}$ phthalocyanine tetrasulfonic acid, and Luxol fast blue MBSN) were found to induce rapid $\text{Ca}^{2+}$ release from actively ($\text{Ca}^{2+}$) loaded SR. Pre-reduction of AB with reducing agents causes loss of stimulating effect indicating that the compounds are acting via redox sensitive interactions with the RyR. (J. J Abramson, Cronin, and G. Salama 1988)

**Quinones:** Naphthoquinone and benzoquinone have been shown to activate channel activity by oxidation of hyperactive cysteine thiols on the RyR1 and RyR2 (Feng et al. 1999).

**Coumarin maleimide 7-diethylamino-3-(4'-maleimidylphenyl)-4-methylcoumarin (CPM):** CPM (1-50 nM) binds to RyR-triadin hyper-reactive thiols when RyR is in a closed state. Covalent interactions of those thiols with nanomolar CPM inhibits gating activity. (G. Liu et al. 1994)

**1,4-Napthoquione:** 1,4-Napthoquione (0.4-40pmol/µM) oxidizes hyperactive thiols on RyR-triadin and stimulates $\text{Ca}^{2+}$ release from SR vesicles. (G. Liu et al. 1994)
3 The Anthocyanidins: Pelargonidin and Delphinidin

3.1 Anthocyanidins

Anthocyanidins are one of the six sub-classes of Flavonoids (itself being a subclass of polyphenols). They are compounds of the plant kingdom that have the general structure of a flavylium cation (2-phenylchromenylum) with a C₆C₃C₆ skeleton and with functionalization occurring at specific locations on the structure (see Figure 3.1). Functional groups are typically referred to by the number of the carbon to which it is attached (eg. The C4' hydroxyl or the 4'-OH). Additionally, the compound is sometimes characterized by its individual aromatic rings A, B, and C. Anthocyanidins are salts and usually are identified along with a chloride counter ion. The counter ion dissociates with the main molecule in solution which leave a positive charge on the compound. This charge allows anthocyanidins to absorb light and exhibit intense colors from the orange-reds to the purples. They provide most of the red to purple colors observed in flowers, leaves, fruits, and
vegetables. They share a common function in the phyto-kingdom as pigments used to attract insects for pollination, cellular signaling molecules, anti-fungal protection, and UV light protection. Specific anthocyanidins are identified by how the base structure is functionalized with hydrogens, hydroxyl, methoxy, and sugar groups and the amount and type of substitution contributes greatly to the color properties of the molecule. There are 23 known anthocyanidins (Øyvind M. Andersen and Markham 2006). Anthocyanidins have poor solubility in aqueous solutions but when sugar groups are attached (anthocyanins), they become more soluble and as such, that is the form that is found most in nature with more than 500 known anthocyanins documented. Although anthocyanins are the more prevalent, the functional properties of those compounds are due to the aglycon form (anthocyanidin) and thus most studies focus on that group of compounds (Estevez and Mosquera 2008; J. B. Harborne, T. J. Mabry, and H. Mabry 1975).

The six anthocyanidins that are considered to be the most important mainly due to their widespread distribution in nature are pelargonidin, cyanidin, delphinidin, petunidin, peonidin, and malvidin. Three of these six are considered to be the principle anthocyanidins due to their minimal substituent groups: delphinidin (Dp), cyanidin, (Cy), and pelargonidin (Pg). These three all have the same hydroxyl functionality on the 3, 5, and 7 carbons of the A and C rings. They differ in the amount of hydroxyl groups on the B-ring. The addition of hydroxyl groups to the B-ring ($R_2=OH$ for pelargonidin, $R_2=R_3=OH$ for cyanidin, and $R_1=R_2=R_3=OH$ for delphinidin) causes a bathochromic (red) shift in their absorption spectrum. They
have absorbance maxima at 530nm, 545nm, and 556nm (respectively) when prepared in acidified ethanol (Cabrita, Fossen, and Øyvind M. Andersen 2000).

3.2 Anthocyanidin Structural Equilibrium

Anthocyanidins are native to the acidic conditions of the plant cell. Under these conditions, the anthocyanidins predominately occur in the ‘flavylium cation’ form as shown in Figure 3.1. Upon placing them in a more alkaline aqueous environment, they become very unstable and their structure is altered in various ways depending mainly on the pH but temperature and light can also be effectors.

At very acidic pH, anthocyanidins exist almost exclusively in the flavylium cationic form (AH\(^+\)) (see Figure 3.2). When subjected to higher pHs (~5), anthocyanidins exhibit three distinct chemical reactions involving proton transfers, hydration, and tautomerizations. Upon a pH jump to ~5, deprotonation immediately occurs on the hydrogen (or hydrogens) of the flavylium cation with the lowest bond disassociation enthalpy (BDE) resulting in the quinoidal base form (A) or the quinoidal base anion form (A\(^-\)). The flavylium cation (AH\(^+\)) and the quinoidal base (A) quickly (<1 sec) come to an equilibrium. On a slower time scale (minutes to hours), AH\(^+\) undergoes hydration to obtain the carbinol pseudobase (B) and/or the carbinol pseudobase anion (B\(^-\)). The rate of these reactions are dependent upon the pH of the environment and on the degree and type of substitutions. Higher pH generally leads to a slower transformation to the carbinol form (Brouillard and Delaporte 1977; Brouillard and Dubois 1977). This will depend on the pK\(_a\)’s of the substituents of the different anthocyanidins. In addition, the
quinoidal base transforms to the colorless carbonyl pseudobase (C) form. For cyanidin (the only kinetic study of a related

Figure 3.2 The forms of pelargonidin in neutral pH solutions
anthocyanidin that I found in literature), the half life of this structural
transformation is reported to be 0.5 days (at pH 2.8) (Iacobucci and Sweeny
1983).
3.3 Pelargonidin

Figure 3.3 Molecular Structure of Pelargonidin

Pelargonidin (Figure 3.4) was first obtained by Willstater and Bolton in 1914 by the hydrolysis of pelargonin of Pelargonium zonale. It has absorbance maxima in the 530nm (in EtOH-HCl). Pelargonidin is found in the skin of the red radish, strawberries, petals of geraniums, and the rose (among others). 22 glycosides of pelargonidin are known (J.B. Harborne 1967). Pelargonidin has hydroxyl (–OH) groups at the 3, 5, 7, and \( \text{para} \) 4’ positions. The structural configurations of pelargonidin are shown in Figure 3.3. When exposed to increasing pH conditions, compounds with a monohydroxylated B ring (pelargonidin) exhibit a gradual bathochromic (red) shift up to pH~6, an extensive red shift from pH~6 to pH~7.3, and a stable \( \lambda_{\text{max}} \) from pH7.3 to pH~10-11 (Cabrita, Fossen, and Oyvind M Andersen 2000). These color changes are caused by structural changes as mentioned in the previous sections. Although the first pK\(_a\) (pK\(_{a1}\)) can be determined experimentally, subsequent deprotonation pK\(_a\)’s (pK\(_{a2}\) etc.) are difficult to determine experimentally due to the simultaneous existence of multiple
forms of the compound in solution. Thus, to predict the pK\textsubscript{a} of subsequent deprotonations of each structural form, quantum mechanical calculations determining the disassociation energies of each hydrogen must be performed. The lowest computed BDEs based on the lowest energy structures are considered the first to deprotonated. In reality, there may be several similarly valued low energy structures and several hydrogens with similar valued BDEs which imply a mixture of compounds in solution at all times. When subjected to a neutral pH environment, pelargonidin rapidly deprotonates and changes from the flavylium cation to its quinoidal base form. The first deprotonation is experimentally been shown to be at pH 5.79. The favored deprotonation site as calculated numerically is the 5-OH group. The predicted pK\textsubscript{a2} for the quinoidal base form is at pH 7.05 and located at the 4’-OH. At physiological pH (~7.4), pelargonidin is probably in an anionic quinoidal base form (see Figure 3.2). The transformation to the anionic quinoidal base is very fast coming with equilibrium in less than 1 second. A second transformation to the carbinol pseudobase has slower kinetics with a half life in the 10 minute range (see result section). The pK\textsubscript{a2} for the carbinol form is predicted to be 8.35 and therefore is probably not anionic at the experimental pH of ~7.1-7.4. The structural change to the chalcone form is on order of 12-24 hours (data for cyanidin, not pelargonidin). The predicted pK\textsubscript{a2} for the chalcone is 6.73 also at the 4’ position and therefore it is predicted to be in an anionic form (Borkowski et al. 2005)
3.4 Delphinidin

Delphinidin (Figure 3.4) was first obtained by hydrolysis of delphin (of the larkspur (Delphinium consolida). It has an absorbance maxima of 556nm (EtOH-HCl) and is responsible for rich purple and dark violet colors of berries. It is found in the skin of eggplant, grapes, and pedals of the hyacinth just to name a few. (J.B. Harborne 1967)

Compounds with a trihydroxylated B-ring (delphinidin) show a gradual bathochromic (red) shift up to pH~6, an extensive red shift from pH6 to pH~7.3, an extensive blue shift from pH7.3 to pH~8.5, and then another red shift at even higher pH's. Due to the same (mixture of forms) difficulties as addressed in the pelargonidin section, the $pK_a$'s for
Figure 3.5 The forms of delphinidin in neutral pH solutions
Delphinidin are also predicted numerically rather than experimentally. Delphinidin’s first deprotonation is at pH 5.30 (experimental) and numerical based deprotonation energies predict it to be the 4’ hydrogen. The predicted pK$_{a2}$ is at 6.73 at C5. At physiological pH, delphinidin is predicted to rapidly deprotonate to the quinoidal base anion form with lost hydrogens on 4’ and 5. The carbinol pseudo base has a predicted pK$_{a2}$ on C4’ at 7.25 and thus may be in a mix of its neutral and anionic forms at physiological pH. The predicted pK$_{a2}$ for the chalcone form is 4.9 also at the 4’ position and therefore it is predicted to be in a neutral form as well (Borkowski et al. 2005).
4 Materials and Methods

4.1 Sarcoplasmic Reticulum Vesicles

Sarcoplasmic reticulum was isolated from fast twitch back and hind leg skeletal muscle extracted from 2-3 kg New Zealand white male rabbits according to the methods of MacLennan et al (1970) with modifications. Live rabbits were euthanized, bled, and put on ice. Removed muscle was also put on ice. Fat and red muscle was trimmed from the fast twitch muscle and discarded. Fast twitch muscle was cut to approximately 1 inch pieces and placed into a cold proteolysis inhibiting buffer of 120mM NaCl, 10mM imidazole, 100µM DTT, 10µM EGTA, and 100µM leupeptin at pH 7.4. Muscle was then homogenized with a common household blender (Waring) by a 4X repeating cycle of 15 sec at low speed, 60 sec high speed, and 30 sec of no blending. Homogenate is then centrifuged at 1,600×g for 10 minutes. Supernate is then filtered through four layers of cheesecloth to remove excess debris and fat. Remaining homogenate is adjusted to pH 7.4 with imidazole and centrifuged at 10,000×g for 15 minutes to remove mitochondria. Supernate is retained and centrifuged at 44,000×g for an additional 70 minutes to collect SR. Supernate is discarded and remaining pellet is homogenized to approximately 10mg/ml with a buffer of 120mM NaCl, 10mM imidazole, 100µM DTT, and 50µM EGTA at pH 7.4. The resulting homogenate is then centrifuged at 7,500×g for 10 min to remove any remaining myosin. Supernate is retained and centrifuged at 78,000×g to obtain the SR pellet. The
pellet is then homogenized to approximately 20mg/ml in a buffer of 100 mM KCl and 20 mM HEPES at pH 7.0. Aliquots of SR are put in cryo-storage tubes and stored in liquid N$_2$ until needed.

The protein concentration of the SR is then quantified by the method of Kalckar (1947). The 280nm absorbance of aromatic amino acids tyrosine and tryptophan combined with the 230nm absorbance of the peptide backbone can be utilized to determine the total protein concentration (see Equation 4.1). Equation 4.2 is a correction factor for the possible contamination of nucleic acids which absorb at 230nm.

**Equation 4.1**

$$[SR]_{mg/ml} = 1.45 \times (Abs_{280}) - 0.74 \times (Abs_{260})$$

**Equation 4.2**

$$[SR]_{mg/ml} = 0.185 \times (Abs_{230}) - 0.075 \times (Abs_{260})$$

### 4.2 Reagents

Ryanodine was purchased from CalBiochem (San Diego, CA), $^3$[H]-Ryanodine was purchased from Du Pont – New England Nuclear Co. (Boston, MA), Pelargonidin chloride (CAS#134-04-3) was purchased from Sigma Aldrich, Delphinidin chloride (CAS#528-53-0) was purchased from Sigma Aldrich and
Extrasynthese (France), Antipyrilazo III and A23187 was purchased from Sigma Aldrich, EtOH was analytical grade 200 proof, anhydrous >99.5%, all water used has a resistivity of 18 Mohm and was obtained from a Mili-Q System (Millipore Co), all other reagents (buffers, salts, etc) were of typical laboratory grade unless noted.

Pelargonidin and delphinidin were prepared at 10mM concentrations in EtOH and stored at ~4°C between experiments.
Stock solution was diluted to lower concentration in EtOH in necessitated by the experiment.

4.3 Methods

4.3.1 Passive Ca$^{2+}$ Loading of SR Vesicles
Ca$^{2+}$ was passively loaded into SR vesicles by incubating undiluted stock SR (~20mg/ml protein) in 0.5mM to 1mM CaCl$_2$ by the procedure as follows. SR was removed from LN$_2$ storage and thawed by rubbing between the hands until no solid SR particles were observed. Multiple storage tubes are thawed and combined as needed to obtain an amount necessary for the experiment. An appropriate amount of 100mM CaCl$_2$ is added to the SR and then vortexed for 3-5 seconds. The tube is then plunged into LN$_2$ and allowed to freeze solid for ~5-
10 seconds. The tube is then thawed again as previously described and placed under crushed ice overnight.
4.3.2 \( \text{Ca}^{2+} \) Efflux Measurements

\( \text{Ca}^{2+} \) release from SR vesicles can be monitored and quantified by spectrophotometric methods utilizing polycationic indicator dyes that change absorbance with corresponding changes in sample free \( \text{Ca}^{2+} \) concentration (Scarpa, Brinley Jr, and Dubyak 1978). The polycationic indicator dye Antipyrylazo III (APIII) and a time sharing dual wavelength spectrophotometer (Chance et al. 1975) (purchased from the University of Pennsylvania) was utilized for all efflux experiments. The spectrophotometer is a rotating wheel type with bandpass filters that pass 720nm and 780nm wavelengths from a halogen source. The passed wavelengths are incident on a photomultiplier tube which is connected to a multiplexing amplification control module. The control module conditions the signals by taking the difference between the (\( \text{Ca}^{2+} \) responsive) 780nm signal from the (\( \text{Ca}^{2+} \) non-responsive) 720nm signal. The difference is then output as an analog voltage signal, digitized, and recorded on a PC (AxoScope software, Axon Instruments (now Molecular Devices)). An APIII buffer containing 100mM KCl, 20mM HEPES, and 200µM APIII at pH 7.4 was used. SR vesicles were loaded with \( \text{Ca}^{2+} \) as previously described. A typical experiment involves 2 ml of the APIII buffer added to a 4.5ml polystyrene cuvette with 1cm path length and a magnetic stir bar. Approximately 40µl-50µl (0.4-0.5 mg/ml final protein concentration) of prepared SR is pipetted into the APIII buffer. The intravascular \( \text{Ca}^{2+} \) concentration is now approximately 50-40 fold greater.
than the extravascular Ca\(^{2+}\). The concentration gradient forces release of Ca\(^{2+}\) through the channel. APIII's spectrum changes in response to the rising Ca\(^{2+}\) concentration and the resulting signal change is recorded for further analysis. At the completion of a Ca\(^{2+}\) efflux experiment, the Ca\(^{2+}\) ionophore A23187 is added (2 µl @ ~1.0 ug/ml) to initiate release of all unreleased intravascular Ca\(^{2+}\) (to determine total vesicle loading capability and to assist in data analysis). A known amount (~5 µM) of CaCl\(_2\) can also be added for Ca\(^{2+}\) signal calibration. To assay for the effect of possible modulators, the modulator is added immediately (<1 sec) prior to the addition of the SR. The resultant signal is compared to the signal of the control. To ensure that the increasing Ca\(^{2+}\) signal is from Ca\(^{2+}\) leaving the RyR1, the potent inhibitor Ruthenium Red is added immediately prior to the SR. Ruthenium Red control should block the release of Ca\(^{2+}\) from the SR vesicles.

To quantify the results, the data representing the Ca\(^{2+}\) release phase is fit to a first order exponential rise to max equation (Sigma Plot). Typically, the initial rate of Ca\(^{2+}\) release is calculated by taking the product of the value of maximum amount of Ca\(^{2+}\) released (µM) and the rate constant (s\(^{-1}\)) which gives the initial rate of Ca\(^{2+}\) in units of µM/s. The initial rate of Ca\(^{2+}\) release is then compared.

It was verified that Pg and Dp do not significantly alter the Ca\(^{2+}\) response of APIII.
4.3.3 Ryanodine Binding

4.3.3.1 Ryanodine

Ryanodine (Figure 4.1) is an alkaloid found in the plant Ryania speciosa of South America. It has a high affinity to the RyR and locks it in a varying degree open state depending on concentration. Due to it's specific affinity to the RyR and it's use as a label for the first purification of the RyR, the RyR is commonly known as the ryanodine receptor.

![Molecular Structure of Ryanodine](image)

Figure 4.1 Molecular Structure of Ryanodine

The rate of ryanodine binding to the RyR depends on the open probability of the RyR. The higher the open probability of the RyR, the greater the rate of Ry
binding. By using tritiated ryanodine, the amount of ryanodine that binds to the
RyR in a specified amount of time can be quantified by a scintillation counter
and be indirectly used to the open state of the RyR. The Beckman LS6500
scintillation counter is utilized for this purpose.

4.3.3.2 Binding Assays
SR (0.5 mg/ml) is combined with a standard ryanodine buffer of 250 mM KCl, 15
mM NaCl, and 20 mM HEPES at pH 7.1. [³H]-ryanodine is added to obtain a final
[³H]-ryanodine concentration of 2 nM. Mixtures are kept on ice. Solutions of
pelargonidin and delphinidin were taken from stock and diluted to obtain a
useable stock concentration if required. [³H]-ryanodine, SR, and pelargonidin or
delphinidin are mixed to obtain desired final concentrations. Mixtures are placed
in a 37°C water bath for the period of time needed for the experiment. At the end
of the time period, the ryanodine binding reaction is then quenched by rapid
filtration through Whatman GF/B glass fiber filter sheets on a Brandel cell
harvester. Sample tubes are rinsed 2x using a standard rinse buffer similar to
standard ryanodine buffer. Filtered samples are placed in scintillation vials with
5 ml scintillation fluid, placed on a shaker overnight, and then counted on a
Beckman L-6500 scintillation counter. RyR non-specific binding is measured by
addition of a 200X excess of non-labeled ryanodine to a control sample. Total
activity is measured by counting an unfiltered sample (2 nM [³H]-ryanodine).

Binding quantification is determined by the following equations:

\[ B_{\text{pmol/mg}} = \frac{1}{[SR_{\text{sample}}] \times vol_{\text{sample}}} \times \frac{dpm_{\text{sample}} - dpm_{\text{NS}}}{SA} \]

**Equation 4.3**

Where

\[ SA = \frac{dpm_{TA}}{[Ry_{\text{total}}] \times vol_{\text{ryanodine}}} \]
5 Results

5.1 Spectrometric Analysis of Ca$^{2+}$ Efflux from SR Vesicles

5.1.1 Pelargonidin Inhibits / Delphinidin Stimulates - Ca$^{2+}$ Efflux from SR Vesicles

This experiment was performed as noted in the methods section. Pg and Dp were added to the APIII buffer immediately (within 1 sec) prior to the addition of the Ca$^{2+}$ loaded SR vesicles. At the concentrations tested (25µM and 50µM), the Pg and Dp stock solutions added 0.25% and 0.5% EtOH (vol) respectively to the APIII buffer. To minimize any effect that the EtOH may have on the RyR and Ca$^{2+}$ efflux, 0.5% EtOH was also added to the control trials. Ruthenium Red (5µm) was utilized to ensure Ca$^{2+}$ efflux was occurring through the RyR and not through other means such as membrane diffusion. Results show (Figure 5.1) that 25µM and 50µM Pg both inhibit the release rate of Ca$^{2+}$ from the RyR in a dose dependent manner and that 25µM and 50µM Dp both stimulate the release rate of Ca$^{2+}$ from the RyR in a dose dependent manner with respect to the control. It is worth noting that 50µM Pg exhibits a $[Ca^{2+}]_{\text{max}}$ that is less than half of the $[Ca^{2+}]_{\text{max}}$ for all other traces. This may be due the existence of a high affinity Pg binding site that is exposed at higher levels of inhibition (higher Pg concentrations). Once the site is revealed, the abundant levels of Pg can bind strongly and effectively close the channel to Ca$^{2+}$ flux. This is highly speculative.
however and similar results were not duplicated in other concentration dependent experiments (see Figure 5.3).
Figure 5.1 Dp stimulates Ca^{2+} release and Pg inhibits Ca^{2+} release from the RyR1 of skeletal SR.

Traces of Ca^{2+} efflux from Ca^{2+} loaded skeletal SR vesicles. Stock SR vesicles (~20 mg/ml protein) incubated with 1 mM CaCl_{2} overnight (~12-16 hours) on ice. Experiments performed in 200µM APIII, 100mM KCl, 20mM HEPES, pH 7.4. Ca^{2+} release is stimulated by dilution of SR into APIII buffer (~40:1). Final protein concentration = 0.3-0.5 mg/ml. Ca^{2+} release is stimulated by both 25µM (▲) and 50µM (△△ △△) Dp and inhibited by both 25µM (▼▼ ▼▼) and 50µM (▽▽ ▽▽) Pg with respect to the control (■■ ■■) (0.5% EtOH). 5µM (●● ●●) Ruthenium Red added to verify release is through RyR1. Both Pg and Dp were added to APIII buffer immediately prior to addition of SR. Error bars represent standard deviation for n=4 for control and n=3 for all other data.
Figure 5.2 Initial Rate of Ca$^{2+}$ Efflux from SR is Modulated by Pg and Dp.

Initial rate of Ca$^{2+}$ release is stimulated by both 25µM and 50µM Dp and inhibited by both 25µM and 50µM Pg with respect to the control (0.5% EtOH). Data from figure 5.1 were fit to first order rise to max exponential functions and results used to calculate the initial rate. Experiments performed in 200µM APIII, 100mM KCl, 20mM HEPES, pH 7.4. Ca$^{2+}$ release is stimulated by dilution of SR into APIII buffer (~40:1). Final protein concentration = 0.3-0.5 mg/ml. Both Pg and Dp were added to APIII buffer immediately prior to addition of SR. Error bars represent standard deviation for n=4 for control and n=3 for all other data.
5.1.2 Pelargonidin Modulates the Initial Rate of Release in a Biphasic Manner

The initial rate of Ca\(^{2+}\) efflux from the SR is stimulated by 1µM and 5µM concentrations of Pg. Concentrations from 40µM to 160µM showed increased inhibition of Ca\(^{2+}\) efflux. A traditional explanation for this type of biphasic behavior is the interaction with more than one class of binding sites on the RyR. At low micromolar concentration of Pg, binding to high affinity at binding sites results in channel activation, while at higher concentrations, binding to a lower affinity site results in inhibition of the channel. This may be the case here but Pg also can show prooxidant behavior at low concentrations and antioxidant behavior at higher concentrations (see chapter 1 and 3) and this characteristic may also contribute to the biphasic character shown here. The data in this section doesn’t support the decreased \([Ca^{2+}]_{\text{max}}\) at higher Pg concentrations (Figure 5.1). The reason for this inconsistency is not known and warrants replicate trials to confirm results. The complex effect that Pg has on the RyR coupled with the lack of more detailed concentration dependence data would render further quantitative analysis as questionable and thus kinetic analysis will not be presented.
Figure 5.3 Pg modulates the initial rate of Ca\(^{2+}\) efflux from skeletal SR in a biphasic manner.

Initial rate of Ca\(^{2+}\) release is stimulated by 1 µM and 5 µM Pg and inhibited by Pg concentrations greater than 10 µM with respect to the control (0.5% EtOH). Experiments performed in 200µM APIII, 100mM KCl, 20mM HEPES, pH 7.4. Ca\(^{2+}\) release is stimulated by dilution of SR into APIII buffer (~40:1). Stock SR vesicles (~20 mg/ml protein) incubated with 1 mM CaCl\(_2\) overnight (~12-16 hours) on ice. Pg was added to APIII buffer immediately prior to addition of SR. Final protein concentration = 0.3-0.5 mg/ml. Results shown are for one trial.
5.1.3 Delphinidin stimulates Ca\textsuperscript{2+} release from the RyR of the SR

Delphinidin stimulates the initial rate of Ca\textsuperscript{2+} efflux from the RyR for all concentrations tested (Figure 5.3). EC\textsubscript{50} = 9.3µM +/- 1.3µM (initial rate data (Table 5.4) fit to a four parameter logistic function (Figure 5.5)). The data is the result of only one trial at each concentration and therefore should be repeated to obtain statistical confidence. The data value at 80µM is approximately 30% less, and not in agreement with, the surrounding data point at 40µM and 160µM. The reason for this is unknown but I suspect that this is an artifact of an experimental error and warrants additional experimental confirmation.
Figure 5.4 Dp stimulates initial rate of Ca$^{2+}$ release from skeletal SR.

Initial rate of Ca$^{2+}$ release is stimulated by Dp in a concentration dependent manner with respect to the control (~0.5% EtOH). Experiments performed in 200µM APIII, 100mM KCl, 20mM HEPES, pH 7.4. Ca$^{2+}$ release is stimulated by dilution of SR into APIII buffer (~40:1). Stock SR vesicles (~20 mg/ml protein) incubated with 1 mM CaCl$_2$ overnight (~12-16 hours) on ice. Dp was added to APIII buffer immediately prior to addition of SR. Final protein concentration = 0.3-0.5 mg/ml. Results shown are for one trial.
Figure 5.5 Dp - Dose Response Curve.

Dose-response data from figure 5.4 fit to a four parameter log function and the half maximal effective concentration (EC$_{50}$) was obtained (Sigma Plot). EC$_{50}$=9.3µM +/-1.3µM. Data point at 80µM (open circle) omitted from curve fit. Experiments performed in 200µM APIII, 100mM KCl, 20mM HEPES, pH 7.4. Ca$^{2+}$ release is stimulated by dilution of SR into APIII buffer (~40:1). Stock SR vesicles (~20 mg/ml protein) incubated with 1 mM CaCl$_2$ overnight (~12-16 hours) on ice. Dp was added to APIII buffer immediately prior to addition of SR. Final protein concentration = 0.3-0.5 mg/ml. Results shown are for one trial.
5.1.4 Pelargonidin and Delphinidin; Different Forms, Different Effects

Both pelargonidin and delphinidin undergo structural changes when exposed to weakly acidic to alkaline solutions (see chapters 1 and 3). The Ca\(^{2+}\) efflux assay has the benefit of being able to more tightly control the time in which Pg and Dp are exposed to the environmental conditions of the assay. In this experiment, stock Pg and Dp were diluted 1:1 in 20mM HEPES at pH7.4 and allowed to set for times periods of 1 hr, 2 hrs, and 4 hrs (all +/- 15 minutes). The samples were then used to determine their effect on Ca\(^{2+}\) efflux. Significant structural changes in Pg and Dp take place in this time period at pH7.4. Results show (Figure 5.6) that Dp loses its ability to stimulate the RyR by a factor of almost 2 after a 1hr exposure to pH7.4. An additional decrease in effectiveness is also seen after 2hrs and 4hrs of dilution time with the final effect being only a slight stimulation of channel activity as compared to the control. Pg loses its inhibitory effect completely after 1 hour of exposure to pH7.4 and its effects on the RyR seem to change from inhibitory effect to a slight stimulating effect after 1hr of dilution.
Figure 5.6 Time of exposure to pH7.4 alters the modulation effects that Dp and Pg have on the initial rate of Ca\textsuperscript{2+} release on skeletal SR.

The inhibition effects of Pg (25µM) are lost after it is exposed to pH 7.4 for 1 hour and slight stimulatory effects are exhibited at longer exposures (2 and 4 hours). Dp (25µM) loses approximately half of its stimulatory effect after 1 hour and continues to lose its stimulatory action at longer exposures (2 and 4 hrs). Experiments performed in 200µM APIII, 100mM KCl, 20mM HEPES, pH 7.4. Ca\textsuperscript{2+} release is stimulated by dilution of SR into APIII buffer (~40:1). Stock SR vesicles (~20 mg/ml protein) incubated with 1 mM CaCl\textsubscript{2} overnight (~12-16 hours) on ice. Dp was added to APIII buffer immediately prior to addition of SR. Final protein concentration = 0.3-0.5 mg/ml. Error bars represent standard deviation for n=3 trial for the control, n=1 for 4hr Dp trial, and n=2 for all other trials.
5.2 Ryanodine Equilibrium Binding

5.2.1 Pelargonidin and Delphinidin Modulate \[^3\text{H}\]-Ryanodine Binding

The equilibrium binding assay is carried out over for 3 hours to ensure chemical equilibrium of the binding of ryanodine to the RyR. Qualitatively, Figure 5.9 shows that both Dp and Pg stimulate equilibrium binding with respect to the control but with Dp stimulating the channel more effectively than Pg. Quantitative analysis and determination of the EC\(_{50}\) would require an assay that includes a broader range of concentrations (greater than 25µM) and thus was not performed. Interestingly, in this assay, Pg exhibits a stimulatory effect. The reason for this is unknown but it can be hypothesized that the change is related to the structural and function changes that Pg undergoes during the 3 hour incubation at neutral pH conditions required for this assay. This stimulatory effect by Dp is consistent with the stimulatory effect seen in the Ca\(^{2+}\) efflux studies.
Figure 5.7 Pg and Dp stimulate ryanodine equilibrium binding.

Both Dp (□) and Pg (○) stimulate ryanodine equilibrium binding in a concentration dependent manner with respect to the control. Samples contain 4nM Ryanodine, 0.5 mg/ml SR vesicles, 50µM Ca\textsuperscript{2+} in pH 7.1 binding buffer. SR vesicles and ryanodine solutions kept on ice prior to dispensing into incubation tubes. Addition sequence: SR (1 mg/ml) in Ry buffer, \textsuperscript{3}H\text{Ry} in Ry buffer, and Pg or Dp. After addition of Pg/Dp samples are put in 37°C bath, removed after 3hrs, and the reaction is quenched by rapid filtration.
5.3 Short Duration Ryanodine Binding

Short duration Ry binding measures Ry binding over a time period of generally less than 20 minutes but other time periods can also be used. Figure 5.8 shows the results of this assay. Delphinidin clearly causes an increase in the initial rate of ryanodine binding with increasing concentration. Attempts made to fit the data to dose response curves did not converge. The lack of strong inhibition in this experiment by Pg can be partially attributed to the low level of binding by the control. In addition, changes in Pg’s structure and the resulting change in its effect on the RyR may also be of importance. In an attempt to enhance the background level of binding and better reveal the effects of Pg on ryanodine binding during initial exposure to pH7.1 buffers, assays were conducted with both 5mM and 10mM caffeine, a known RyR stimulator, added to the experiment. Figure 5.9 shows that caffeine stimulated binding is strongly inhibited by Pg.
Figure 5.8 Pelargonidin inhibits caffeine stimulated ryanodine binding

Pelargonidin inhibits 5mM (○) and 10mM (□) caffeine stimulated ryanodine binding in a concentration dependent manner with respect to the control. Samples contain 4nM $^3$H-ryanodine, 0.5 mg/ml SR vesicles, 50µM Ca$^{2+}$ in pH 7.1 binding buffer. SR vesicles and ryanodine solutions kept on ice prior to dispensing into incubation tubes. Addition sequence: SR (1 mg/ml) in Ry buffer, $^3$H-Ry in Ry buffer, and Pg. After addition of Pg, samples are put in 37°C bath for 5min and then removed and the reaction quenched by rapid filtration.
Figure 5.9 Comparison of $^3$H-ryanodine binding effects of Pg at short and long exposures to a pH 7.1 aqueous buffer.

When Pg's exposure to a pH 7.1 buffer is limited to 5 (▼) or 6 (▽) minutes, Pg inhibits $^3$H-ryanodine binding to the skeletal RyR. When the structural changes are allowed to proceed for 3 hours (▲), Pg stimulates $^3$H-ryanodine binding to the skeletal RyR. Samples contain 4nM $^3$H-ryanodine, 0.5 mg/ml SR vesicles, 50µM Ca^{2+} in pH 7.1 binding buffer. SR vesicles and ryanodine solutions kept on ice prior to dispensing into incubation tubes. Addition sequence: SR (1 mg/ml) in Ry buffer, $^3$H-Ry in Ry buffer, and Pg. After addition of Pg, samples are put in 37°C bath for 5min and then removed and the reaction quenched by rapid filtration.
5.4 Spectrophotometric Characteristics of Pelargonidin and Delphinidin

5.4.1 Absorbance Spectrum of Pg and Dp

The absorbance spectrum exhibited by Pg and Dp is very dependent upon the pH of the solution as well as the time that Pg and Dp have been exposed to that pH. Figures 5.11 and 5.12 show the spectrums of Pg and Dp respectively at various pH’s. The spectrum was taken as quickly as possible (~3sec) after the dilution of 40 µM (final concentration) Pg or Dp from its EtOH stock solution into 1mM Tris buffer at various pH’s. Results show the characteristic bathochromatic shifts of their spectrums that are expected at increasing pH (Cabrita, Fossen, and Øyvind M. Andersen 2000).
Figure 5.10 pH dependent absorbance spectrum of pelargonidin

40uM Pg pipetted from 10mM stock in EtOH to 1mM TRIS (pH’d with HCl and/or KOH) and spectrum collected within 5 seconds. Solid line: pH=6.5, long dashes: pH=7.0, short dashes: pH=7.5, dotted trace: pH=8.0
Figure 5.11 pH dependent absorbance spectrum of delphinidin

40uM Dp pipetted from 10mM stock in EtOH to 1mM TRIS (pH’d with HCl and/or KOH) and spectrum collected within 5 seconds. Solid line: pH=6.5, long dashes: pH=7.0, short dashes: pH=7.5, dotted trace: pH=8.0
5.4.2 pH Dependent Transformation Kinetics of Pg and Dp

Upon dilution into a neutral pH solution, Pg and Dp rapidly deprotonate into their respective quinoidal base forms. This transformation occurs in a time frame faster than that required to add, mix, and collect the initial spectrum and thus cannot be followed using the technique described here. The subsequent slower transformation from the quinoidal base form to the carbinol pseudobase form is shown in Figure 5.13. The time required to add, mix, and collect the initial spectrum is on the order of a few seconds and therefore this measurement will only provide an estimate of the actual kinetics. However, this data is in agreement with the literature in that the transformation kinetics of these compounds increase with increasing acidity. In addition, the relative concentration of the quinoidal form with respect to the (colorless) carbinol form is dependent on pH. Unfortunately, the equilibrium constants for these compounds are not known and concentrations of the different forms cannot be determined. The kinetics of Pg and Dp are similar and when fitted to a first order decay equation, the $t_{1/2}$ is on the order of 100 seconds at pH 7.0. The kinetics significantly slow down at pH 7.5 and it is therefore not possible to estimate the $t_{1/2}$ at this pH.
Figure 5.12 pH dependent transformation kinetics of Pg and Dp

Kinetics of the decrease in absorbance of the quinoidal form of Pg and Dp over 300 seconds. Traces for pH 6.5 (t₁/₂~50 sec), 7.0 (t₁/₂~50 sec), 7.5 (t₁/₂~not enough data) and 8.0 (t₁/₂~not enough data) are shown. Black traces are Dp and Lt. Grey traces are Pg. Solid line: pH=6.5, long dashes: pH=7.0, short dashes: pH=7.5, dotted trace: pH=8.0.
6 Discussion

The complex nature of pelargonidin and delphinidin with respect to their pH dependent molecular structures and properties makes it difficult to quantitatively assess the two compounds with regards to RyR modulation. However, calcium efflux experiments do allow for assessment of the effects of pelargonidin and delphinidin within seconds of exposure when concentrations of specific forms are known to dominate. Upon exposure to physiological pH solutions, pelargonidin and delphinidin rapidly (~1 second) deprotonate to their corresponding quinoidal base and quinoidal base anion forms whereas the subsequent transformation to the carbinol pseudobase is on the order of minutes at pH 7 and significantly longer in more alkaline solutions. The Ca$^{2+}$ efflux assays utilized in this thesis can probe the modulatory effect that a compound has on the channel in time periods as short as several seconds and therefore can effectively assay pelargonidin and delphinidin in their quinoidal forms. Additionally, the transformation of pelargonidin and delphinidin to the chalcone form is likely on the order of twelve to twenty four hours (based on similar compounds found in literature) and therefore there is a time period after the transformation into the carbinol pseudobase that the Ca$^{2+}$ efflux assay can probe the effects of that form as well.

Delphinidin is predicted to be in a quinoidal base anion form almost immediately after exposure to a pH 7.4 solution. Hydroxyl groups at positions 4’ and 5 are the predicted deprotonation sites. Calcium efflux results show that delphinidin is a
good RyR1 stimulator with an EC$_{50}$ of 9.3µM. However this result is obtained within the first fraction of a minute after Dp is exposed to the pH 7.4 of the Ca$^{2+}$ efflux buffer. When Dp is allowed to transform into the carbinol pseudobase form, a majority of its stimulatory effects are essentially lost. This implies that the quinoidal base form is primarily responsible for Dp's stimulatory effects on the RyR. This also implies that the carbinol pseudobase form doesn't have a strong modulatory effect. Although there is still a small stimulatory effect seen even after the carbinol is formed, this may be due to the fact that there is still an equilibrium concentration (not known) of quinoidal base in solution or that stimulatory autoxidation products of Dp are present in the solution (both superoxide and hydrogen peroxide have been shown to stimulate the RyR1 (T. G Favero, A. C Zable, and J. J Abramson 1995; Kawakami and Okabe 1998)). $[^{3}\text{H}]-$ryanodine binding experiments at both short $[^{3}\text{H}]-$ryanodine incubation times (~ 6 minutes (data not shown)) and long incubation times (~ 3 hours) also show stimulatory effects. Again, the stimulatory effects for longer time periods are suspect in part due to the autoxidation capabilities of Dp.

With pelargonidin, the modulating mechanism is more complex. Pelargonidin is also predicted to be in a quinoidal base anion form immediately after exposure to a pH 7.4 solution. Hydroxyl groups at positions 5 and 4' are the predicted deprotonation sites but the 5 and 3 are close in energy and probably co-exists in significant concentrations. Calcium efflux results imply that the quinoidal form of pelargonidin is a RyR1 inhibitor. In contrast, the carbinol pseudobase does not
seem to be effective at inhibiting the RyR. There is even a slight hint of some channel stimulatory effects. \(^{3}\text{H}\)-ryanodine binding experiments performed with short \(^{3}\text{H}\)-ryanodine incubation times (~ 5 minutes) also show activation of receptor binding. As is the case with delphinidin, the structural form 3 hours after exposure to neutral pH conditions is also the carbinol pseudobase although, for pelargonidin, the predicted (lowest) pK\(_a\) of that form is 8.32 (3-OH) and therefore it probably exists with a neutral charge. Longer \(^{3}\text{H}\)-ryanodine incubation times (~ 3 hours) exhibited significant channel stimulatory effects on the RyR. Unlike Dp, Pg is not known to autoxidize and therefore the generation of oxidation products doesn’t seem to be a probable explanation for the stimulatory action.

For the quinoidal base forms, both compounds are predicted to be planar and both compounds are charged with the deprotonation sites on the 5 and 4’ oxygens. However, the relative order of the acidities of the hydrogens of those groups are reversed. Dp loses its first hydrogen at O4’ (experimental pK\(_{a1}\)~5.30) and its second hydrogen at O5 (predicted pK\(_{a2}\)~6.30) whereas pelargonidin loses its first hydrogen at O5 (experimental pK\(_{a1}\)~5.79) and its second hydrogen at O4’ (predicted pK\(_{a2}\)~7.05). The second predicted deprotonation of pelargonidin is also close to the pH of the assays performed in this thesis (pH7.1 for ryanodine binding and pH7.4 for Ca\(^{2+}\) efflux) which means that pelargonidin will exist as more of a mixture of both the neutral quinoidal base and the anionic quinoidal base forms. This may impart more of a hydrogen donor characteristic to
pelargonidin relative to delphinidin. The relative order of the deprotonation sites may also impart differing molecular polarization characteristics to Pg. How this may affect the RyR is unknown due to the lack of knowledge regarding the Pg (and Dp) binding site(s). Structural information regarding the carbinol pseudobase forms of these compounds was not found in literature however the predicted $pK_{a2}$ of the carbinol pseudobase forms of pelargonidin and delphinidin are 8.35 (at the 3-OH) and 7.25 (at the 4’ OH) respectively. Thus, when the carbinol pseudobase structures are formed (in neutral pH conditions), pelargonidin carries mostly a neutral charge whereas Dp is in a mixture of neutral and anionic forms (Borkowski et al. 2005).

In conclusion, the distinct and powerful effects that the anthocyanidin compounds studied in this thesis potentially have on skeletal ryanodine receptor mediated Ca$^{2+}$ homeostasis is further proof of the necessity to fully understand the complete effects of nutraceuticals before dietary intake recommendations can be made.
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