History of atrial fibrillation 1628–1819
Harvey – de Senac – Laënnec

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SUMMARY For two centuries after Harvey the arterial pulse was often regarded as independent of the heart beat. This misunderstanding was mainly because of the frequent failure of the irregular heart to elicit a radial pulse: the “pulse deficit” of later times.

Harvey observed ineffective palpitation of the atrium just before death. This was probably atrial fibrillation. He established the origin of the heart beat in the right atrium.

Harvey’s observations were confirmed and extended by de Senac in the mid-eighteenth century. He correlated gross irregularities (palpitation) with necropsy observation of mitral valve disease and dilatation of the left ventricle. He emphasised the origin of the heart’s irregularity from the distended atrium consequent on distension or reflux of blood irritating the atrial wall.

He also commented on disconcerted action and rippling of the ventricular wall before final cessation of movement in a dying heart (ventricular fibrillation).

De Senac’s ideas were a century and a half ahead of his time.

The problem of the pulse
In Chapter 3 of De motu cordis (Franklin’s translation, 1957)¹ are laid out the logical reasons by which Harvey comes to the conclusion:

there is just the one cause of arterial pulsation throughout the body, and that is contraction of the left ventricle. The pulsation of the artery-like vein (pulmonary artery) is similarly related to the contraction of the right ventricle . . . . In correspondence with the tension of the heart the arterial pulsations become larger, more forcible, frequent, rapid, while preserving the rhythm, volume and order of the heartbeat . . . . For the pulse of the arteries is nothing save the impulse of the blood entering these vessels.

This clear understanding had to wait nearly 200 years for further application in the precise study of cardiology. It was debated, and often misunderstood by practitioners of medicine, and indeed Abercromby (1685) in a book on the pulse,² said: “Its source is as mysterious as the source of the Nile”.

The belief, held by many, that arteries had an inherent movement of their own was even supported by Laënnec (1819). In discussing “intermission of the pulsation of the heart”, he noted variations in the heart’s contractions with or without a palpable pulse and commented as follows:

Many considerations prove that the mere examination of the pulse is insufficient to inform us of the state of the circulation and must often lead us into error . . . .

After what has been said and after its general uncertainty avowed by the most experienced practitioners it may seem surprising that the practice of feeling the pulse has been so generally followed in all ages. The reason for this practice is, however, sufficiently obvious. It is easy of performance and gives little inconvenience either to the physician or to the patient . . . . The facts above stated relative to the discordance existing between the pulsation of the heart and of the arteries, more especially as to strength, are contrary to the more general opinion of modern physiologists who consider the action of the arteries as entirely dependent on that of the heart. Bichat himself has fallen into this error.

(Laënnec, Forbes’ translation, 1821).³

Marie François Bichat (1771–1802) was the distinguished anatomist of the Hôtel Dieu, whose observations were much quoted by Laënnec. Bichat promoted the study of tissues and membranes before the era of the microscope. Thus the “pulse deficit” of atrial fibrillation seems to have contributed to the centuries long failure of the profession to accept Harvey’s firm demonstration that the beating heart created the pulse. Sometimes the beat was too weak to be felt at the radial artery.

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Origin of the heart beat

Harvey's meticulous observations (chapter 4, Franklin's translation1) had already established that the heart beat was initiated in the atria, and the right atrium in particular. From his many direct observations on the heart in animals, fishes, frogs, eels, birds, and domestic animals, Harvey gives the following account:

The movement is seen to begin from the auricles and to pass on to the ventricles. With everything becoming more sluggish as the heart lies dying, and also in fishes and the colder bled animals, these two movements become separated by an interval of inactivity . . . . Finally as it sinks to death (the ventricle) ceases to reply with a proper movement to the pulsating auricle. The heart (i.e. ventricle) ceases to beat before the auricles so that the auricles may be said to outlive it . . . . With all the other parts inactive and dead the right auricle goes on beating so that life appears to linger longest in this auricle . . . . If at this time you cut the apex of the heart with a pair of scissors you will see blood flow out from the wound with each beat of the auricle. You will thus realize that the blood gets into the ventricle not through any pull exerted by the distended heart but through the driving force exerted by the beat of the auricles.

Harvey continued:
But I . . . have noticed, that after the heart proper, and even the right auricle were ceasing to beat and appeared on the point of death, an obscure movement, undulation/palpitation had clearly continued in the right auricular blood itself for as long as the blood was perceptibly imbued with warmth and spirit.

Harvey must thus be credited with the first direct observation of a fibrillating atrium.

Richard Lower (1631–1691) began as research assistant to Thomas Willis and was introduced into the Royal Society by Boyle. He was the best English physiologist after Harvey and later practised medicine in London. He showed that blood changed colour to bright red in passage through the lungs and that asphyxia prevented this. He laid down in his Tractatus de Corde (1669, p. 75),4 “The heart is definitely a muscle and has movement exactly similar to other muscles. It receives spirits from the abundance of nerves which are inserted into it”.

Vital spirits from the nervous system were supposed to animate the nerves. The idea of nervous origin of the heart beat was to persist for another two centuries.

De Senac's critical analysis

In the mid eighteenth century a new dimension of analysis was added by de Senac (Fig) whose views were much in advance of his time. He built his physiological ideas on the basis of Harvey and Lower, repeating their experiments and applying critical analytical thought.

Jean-Baptiste de Senac (1693–1770), MD Montpellier, became physician to Louis XV and also became a Councillor of State. In 1749 his Traité de la Structure du Cœur de son action et de ses maladies (S) was published and a second edition appeared in 1774.5 This is the first textbook on cardiology. Volume I contains the most detailed account of the anatomy of the heart in adult and in fetal life and the problems of its physiology are considered thoroughly in Volume II. He also published a more concise Traité des Maladies du Coeur (M) in two editions (1774–1783). It is from the second edition and second volume of S and M (“structure” and “maladies”) that the following extracts are made. The posthumous editions were guided through the press by Baron Antoine Portal, but the text had been revised by the author.

De Senac confirmed Harvey (S, p. 149) that in the heart of a dog dying after severing an artery, beating persists longest in the right atrium and also in the termination of the vena cava. Thus the ultimum moriens part of the heart was confirmed in the sinoatrial region.

Although de Senac built on the ideas of Richard Lower that the heart was innervated like other muscles, he asked many difficult questions: why can the heart contract after death? (S, p. 102). Cut the nerves and the heart does not stop? de Senac established irritability of the heart. Touching the heart may make it contract even more strongly. It will contract in response to internal pressure: compare the intestines. He was mystified by the visible contraction of the non-innervated embryo heart of the chick, which he thought might be the result of some still unknown agency. The debate about nervous control of the heart beat continued for another century and a half.

Just as Harvey had seen fibrillation of the atria in a dying heart, de Senac gave a description of terminal fibrillation of the ventricles (S, p. 151):

Before these parts (of the heart) lose their action the harmony of their movements is disconcerted: there is neither order nor regular succession . . . . When their (muscle) mass no longer makes any effort their fibres are still agitated. They form oblique ripples and there is seen, according to Pechlin, an undulation of many varieties moving from the base to the apex in a spiral form following no doubt the direction of the superficial fibres.

Johannis Nicolai Pechlin (1646–1706) observed death in an open chest wound (Historia Vulneris Thoracici, 16836). Born in Holland, he practised in Kiel.
Clinico-pathological correlations

When he turned his attention to clinical abnormalities of the heart, de Senac was linking his observations with pathological anatomy in the same way as Morgagni, his slightly older contemporary.

Giovanni Battista Morgagni (1682–1771), was Professor of Theoretical Medicine at Padua from 1720 until his death. He published his book on *The Seats and Causes of Diseases investigated by Anatomy* in 1761. This established a pattern of observation by which the clinical manifestations of disease might be better understood.

De Senac devoted much attention to palpitation
and his necropsy correlations are best described in his
_Traité des Maladies du Coeur._ He emphasised mitral
valve disease in the causation of palpitation. His mor-
bid anatomical observations are precise. Under "The
particular and proximate causes of palpitation are in
the heart or in the vessels":

Cardiac causes are disorders of its very substance
or of the valves. In one word obstacles which retain
the blood in the ventricles and which under con-
tinuous pressure irritate the heart fibres causing
stronger contractions. Reflux of blood into the
auricles causes contractions of greater force, neces-
sarily precipitating palpitations. When the obsta-
cles in the heart obstruct the entrance to the ven-
tricles, blood accumulates in the cavities of the
auricles and irritates the fibres of their walls. Their
contractions become more lively. (_M_, p. 7–9).

De Senac quotes a case with irregularity of the heart
where the tricuspid valve was “narrowed” in the same
manner as that described by Vieuxsens in the mitral
valve (_M_, p. 9). He gives another description of nec-
ropsy in which “the tricuspid valve was narrowed and
cartilaginous” and “the mitral valve was narrowed,
shut down to the inner surface of the ventricle and
adherent to its walls. The chordae tendineae were
shortened, rendering the cusps immobile” (_M_, p. 10).
This is a precise description of a “funnel” stenosis and
regurgitation of the mitral valve. Then he goes on:

Dilatation of the ventricles may also cause palpita-
tions. This results from obstruction of the exit of
blood which distends the ventricles which thus
contract strongly causing more violent reflux into
the auricles. (_M_, p. 11).

These repeated references to distension of the atria
and the precipitation of contractions by pressure of
the accumulating blood were somewhat prophetic of
the ideas developing in the early nineteenth century
when the irregular pulse of atrial fibrillation was usu-
ally regarded as a diagnostic sign of mitral stenosis,
which was indeed its commonest cause. De Senac’s
observations had, however, also related the irregular-
ity to enlargement of the ventricles as well. But even
here he comes back to the effect of ventricular dilata-
tion on reflux of blood into the atria.

In his discussion of palpitation in general, de
Senac’s remarks are full of observational clinical wis-
dom:

Palpitations consist of beats which are either more
active and more frequent, such as occur in fever or
in any action demanding effort, and even mental
agitation. But these are not illnesses by themselves
and do not leave any sequential damage. When the
body settles down to rest the heart resumes its
normal beat. (_M_, p. 2).

He concluded that the nerves of the heart were impor-
tant in the precipitation of palpitation:

There are, however, palpitations which are lively,
and irregular beats agitate the heart. The com-
plaint is produced by various fortuitous events and
often appearing without any obvious cause, settle-
ling down with rest, but possibly recurring. We
only define as palpitations beats which are lively,
irregular, agitate the heart and form a veritable
malady of the beat of the heart... continuing
during rest... (_M_, p. 2).

When the palpitations are irregular they form a
veritable malady of the beat of the heart acting on
the great arteries. The primary cause of the disor-
der is an irritation... of the tissues of the heart
and a reflux of blood which is directed against the
auricles with more violence... The irritation is
followed by contraction of the muscle and the grea-
ter the force of the ventricles the more the
reflux... (_M_, p. 5).

The reflux agitates the heart or disturbs its
action. (_M_, p. 6).

If the auricles are strained and increased in vol-
ume they cause palpitations... On exercise too
much blood arrives at the sinus of the vena ca-
v... its walls are irritated by the overload of
blood which is driven more forcibly to the right
ventricle and its action becomes more lively: hence
a reflux which is still more violent and beats in
response to this reflux. One sees the same effects
when the auricles are abnormally dilated... their
walls are more irritable and they act more actively
on the ventricles. (_M_, p. 12–13).

In the case of the funnel valve described above,
The ventricles send a violent reflux into the auricu-
les. Distension of the walls of the heart makes
palpitations inevitable. (_M_, p. 11).

In dealing with palpitation he points out that
irregularities can be precipitated by distension of the
stomach, which could be relieved by the use of vari-
ous carminatives, and he noticed that the remedy
whose effects have appeared to be the most constant
and the most prompt in many cases is quinine mixed
with a little rhubarb. “Long and rebellious palpita-
tions have ceded to this febrifuge, seconded with a
light purgative”. This is quoted by Willius and Keys8
from his first edition of 1749, and is prophetic of the
current rational use of quinine first observed and
noted by an astute physician. De Senac’s other deep
insights have had little mention in medical historical
texts.

De Senac’s remarkable insights into the correla-
tions of mitral disease with gross pulse irregularities
were based on his own observations on cardiac muscle
irritability and the atrial origin of the heart beat as
established by William Harvey and expanded by de
Senac’s own meticulous observations. He stood firm
on the atrial origin of the heart beat as established by
William Harvey. Like Harvey he also put much
emphasis on the development of rippling movements
in the dying chambers of the heart atria and also ven-
tricles. These are probably the earliest experimental
observations of fibrillation. He commented, however
(S, p. 52), that “the causes of palpitation are not the causes of the natural heart-beat”, thereby hinting at their ectopic origin.

De Senac advanced understanding to its very limits in his day in exemplary observation and deep critical thought. He also commented on the speculative writings of others: “Rien n'est plus rare que l'exactitude d'observation” (M, p. 8).

Note: The author is responsible for the selection of quotations and translation.

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References

5 de Senac JB. Traité de la structure du cœur, de son action et de ses maladies. 2nd ed., vol. II. Paris: Mequignon l'ainé; 1783.

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