Introduction

Scombroid Syndrome (SS, also known as Scombrotoxism or Scombroid Ichthyotoxicosis) is a recently described pathology due to the consumption of altered fish flesh, in the absence of organoleptic abnormalities [1]. It is a particular type of Kounis Syndrome, a disease in which a cardiovascular event (a coronary vasospasm, a myocardial infarction or a stent thrombosis) occurs as the consequence of an allergic or hypersensitivity, anaphylactic or anaphylactoid insult [2]. This inflammatory basis has a first actor in mast cell [3,4]. Mast cell interacts and activates directly or indirectly other inflammation cells (macrophages, T lymphocytes) [4]. In hypersensitivity mast cells degranulation put in circulation several inflammatory mediators like histamine [4]. Histamine is a biogenic amine that has a major role in the inflammatory and allergic response [6]. Its effects include coronary vasoconstriction, induction of tissue factor expression, and platelets activation [6-8]. These effects justify the name of ‘Histamine fish poisoning’ with which SS is also known. Other inflammation mediators have a cardiovascular effect [2]: chemokines, enzymes such as the neutral proteases chymase, tryptase and cathepsin-D, cytokines, peptides, proteoglycans, growth factors and arachidonic-acid products such as leukotrienes, thromboxane, prostacyclin, PAF, and tumor necrosis factor. All the neutral proteases can activate metalloproteinases which degrade the collagen cap of the plaque, bringing to erosion and then rupture. Cathepsin-D and Chymase have a role in converting Angiotensin I in Angiotensin II, with a vasoconstrictor effect [9]. Vasoconstriction is induced also by leukotrienes [10]. Thromboxane has both a vasoconstrictor and a platelet aggregation mediator effect. All these mediators assume a particular relevance for patients with cardiovascular risk factors, especially for those with a metabolic syndrome [11]. They play also a role in the more general context of the Kounis Syndrome with Histamine playing a central role in SS [2,12].

We performed a systematic research on Pubmed using the key words ‘Scombroid Syndrome’ and ‘Kounis Syndrome’. We found 251 articles but we selected only 23 articles of interest.

SS physiopathology is explained by the high level of Histamine in the altered fish flesh [13]. It is important to underline that freshly caught fish does not contain a sufficient quantity of histamine to determine symptoms [14,15]. The potential risk is linked to fish species (especially the Family of Scombridae, migratory pelagic species such as mackerel, tuna and tuna yellow-fin and non-migratory species such as menhaden and minnows) in which there is a high level of the amino acid histidine in its free form, physiologically present as a buffer system in their tissues, protecting them from the sudden increase of lactic acid [16-19]. A good maintenance of the cold chain prevents the decaboxylation by histidine decarboxylase, an enzyme present in some bacterial species, in histamine [20]. This process occurs typically in early stages of fish deterioration, in the absence of organoleptic abnormalities, when after fish death, there is no more defense against bacterial growth. An interesting hypothesis suggest that some substances present in fish, increase histamine toxicity in human, promote its absorption, or inhibit its inactivation by histamine N-methyltransferase and diamine oxidase [3,21,22]. There are some enhancers in the fish tissues (other amines like putrescine, cadaverine, tyramine, agmatine) which could play a role in the development of
cause massive mast-cell degranulation. Paracetamol is not
be exercised with opioids, which, in Kounis syndrome, may
concentration than the subcutaneous one). Caution must
be administered a sulfite-free epinephrine intramuscularly
(this route has faster onset of action with a more stable
receptors. Epinephrine in Kounis syndrome (typically the
coronary spasm due to unopposed activity of α-adrenergic
but the administration of β-blockers may exaggerate
vasospasm, especially in Type II variant,
Nitrates, calcium channel blockers can be administrated
steroids with resolution of hives in the Type I [28-30].
In this setting some improvement in symptoms has been
reported with antihistamines treatment and possibly with
histological demonstration the presence of eosinophils and
mast cells). The evolution from SS to Kounis Syndrome is a
potentially risk for the patient life that the physician should
remember.

Particular attention should be paid to those patients who
arrive to the Emergency department with cardiovascular
risk factors (arterial hypertension, diabetes mellitus,
dyslipidemia, metabolic syndrome, previous myocardial
infarction): their danger threshold is lower. According
with Athyros et al [25], a new cardiovascular risk disease
estimation could be helpful to better define an optimal
management protocol and pharmacological approach. In
fact several recent evidences justify higher level of alert
in patients with a chronic activation of the inflammatory
cascade, also in those with a not primitively heart disease
[26]. The reason for this kind of approach is attributable
on the central role of endothelium dysfunction between
inflammatory mediators and atherosclerosis [27].

In this setting some improvement in symptoms has been
reported with antihistamines treatment and possibly with
steroids with resolution of hives in the Type I [28-30].
Nitrates, calcium channel blockers can be administrated
to reduce the vasospasm, especially in Type II variant,
but the administration of β-blockers may exaggerate
coronary spasm due to unopposed activity of α-adrenergic
receptors. Epinephrine in Kounis syndrome (typically the
drug of choice in anaphylaxis) may aggravate ischemia and
worsen coronary vasospasm, so only in severe cases should
be administered a sulfite-free epinephrine intramuscularly
(this route has faster onset of action with a more stable
concentration than the subcutaneous one). Caution must
be exercised with opioids, which, in Kounis syndrome, may
cause massive mast-cell degranulation. Paracetamol is not
recommended, especially by intravenous administration,
because it might cause severe hypotension. In Type III
variant is a protocol for myocardial infarction should be
started, with urgent aspiration of intra stent thrombus
[31-33]. There are emerging evidences that the flavonoid quercetin has a good efficacy in blocking mast
cell cytokines in humans [34]. This interesting option
could be effective in the prevention of Kounis syndrome as
well as a diet containing natural flavonoids with mast cell
inhibitors [35].

SS definitive diagnosis implies that allergic symptoms are
present, an antihistaminic therapy is effective and the
presence of high levels of histamine in the fishery product.

SS often does not show itself in a dangerous way but the
physician of the Emergency Care unit should keep in mind
that its evolution is not ever predictable and there is a
potential risk of life if heart is involved, especially in
atopic subjects and in those who develop an anaphylactic
reaction with pre-existing cardiovascular risk factors.
Recognize SS and its development in a Kounis Syndrome can
save life to the patient. In this setting a new definition of the
cardiovascular risk disease, with a combination of the
emerging evidences about the role of quercetin and of
the natural flavonoids, may allow physicians to take early
therapeutic decisions.

Abbreviations
SS-Scombroid Syndrome

Keywords: Acute Coronary Events; Vasospastic
Allergic Angina; Allergic Myocardial Infarction; Allergic
Stent Thrombosis; Scombroid Syndrome; Kounis Syn-

Declaration of Interest
The authors declared no potential conflicts of interest with
respect to the research, authorship, and/or publication of
this article.

References
1. Arnold SH, Brown WD. Histamine (?) toxicity from fish
2. Nicholas G. Kounis, Coronary Hypersensitivity Disorder:
3. Kathleen J. Motil, Nevin S. Scrimshaw. The role of
exogenous histamine in scombroid poisoning. Toxicol Lett.
5. Doherty TM. T cell regulation of macrophage function.