Ricin: an ancient story for a timeless plant toxin

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Abstract: The castor plant (Ricinus communis L.) has been known since time immemorial in traditional medicine in the pharmacopoeia of Mediterranean and eastern ancient cultures. Moreover, it is still used in folk medicine worldwide. Castor bean has been mainly recommended as anti-inflammatory, anthelmintic, anti-bacterial, laxative, abortifacient, for wounds, ulcers, and many other indications. Many cases of human intoxication occurred accidentally or voluntarily with the ingestion of castor seeds or derivatives. Ricinus toxicity depends on several molecules, among them the most important is ricin, a protein belonging to the family of ribosome-inactivating proteins. Ricin is the most studied of this category of proteins and it is also known to the general public, having been used for biocrimes in several cases. Here, the main steps of ricin research are reported with particular regards to its enzymatic activity, structure and cytotoxicity. Moreover, we discuss ricin toxicity for animals and humans, as well as the relation amongst bioterrorism and ricin and its impact on environmental toxicity. Ricin has also been of great utility to develop a number of immunotoxins specific for the elimination of unwanted cells, mainly cancer cells; some of these immunotoxins gave promising results also in clinical trials.

Keywords: castor bean; cancer therapy; immunotoxins; plant toxins; ribosome-inactivating proteins; ricin; rRNA N-glycosylase activity; traditional medicine; folk medicine; bioterrorism

Key Contribution: This manuscript points out the most known plant toxin: ricin. Starting from the use of Ricinus plant in traditional and folk medicine; we highlight the milestones of research on ricin; with particular regards to its enzymatic activity; structure; cytotoxicity; toxicity for animals and humans and the double face of its employ, for biocrimes and medicine.

1. Castor bean in traditional and folk medicine

The toxin ricin derives from Ricinus communis L. (Euphorbiaceae family), also known as castor bean or palma Christi. The genus Ricinus has only one known species: the castor oil plant. The plant possibly originates from Africa and Asia and now is widespread throughout temperate, subtropical and tropical areas, growing as an invasive plant or being cultivated for different purposes also in Westland.

The castor plant has been known since time immemorial and its use in prehistoric era is evidenced by archaeological findings such as that of the Border Cave in South Africa. Traces of wax containing ricinoleic and ricinelaidic acids were found on a thin wooden stick, which was suggested to be a poison applicator, dating back to about 24,000 years ago [1]. The castor seeds and other parts of the castor plant were certainly utilized in ancient Egypt for pharmacological purposes. In the Ebers Papyrus, an Egyptian medical treatise dating back to before 1500 BCE, an entire chapter is dedicated to castor bean that is indicated as abortifacient, laxative, for treatment of abscessual illness, for...
baldness and so on [2]. In the Hearst Papyrus, written approximately in the same period, various
castor plant parts are included as ingredients in some prescriptions for internal use with the aim of
expelling fluid accumulation or promoting diuresis, as well as for external use as poultices for
bandaging [3]. Ancient Egyptians knew the toxicity of castor bean and only small amounts of the
seeds pulp were included in drug preparations for oral ingestion, which were prescribed mostly with
abortifacient or laxative purposes. In addition, a castor seed-containing concoction was
recommended to cure the urinary disease of a possibly diabetic child [4]. Around 400-year BCE, the
father of western medicine Hippocrates prescribed castor bean oil for laxative and detoxifying action
[5]. The Greek herbalist and physician Pedanius Dioscorides (40–90 CE) in De Materia Medica wrote
that castor seeds could be used as expectorant, diuretic, emetic, laxative, anti-inflammatory, to cure
erysipelas, burns, varicose veins, etc [6]. In the same period, Pliny the Elder (23-79 CE) wrote Naturalis
historia, comprising the whole area of antique knowledge. In this encyclopedic work there is a place
also for castor bean [7].

Castor bean was used also in the pharmacopeia of eastern ancient cultures. In Chinese traditional
medicine, castor seeds were recommended for their anthelmintic activity; seed poultice and leaf juice
were prescribed for external use to treat ulcers and chronic wounds, whereas the latex was instilled
in the ear to recover patient from rhinitis (reviewed in [8]). Castor plant is called erandah in Sanskrit,
because of its reputation of being a remedy for all kind of diseases. In Ayurveda, castor plant is used
for rheumatic affections, as well as for gastropathy, constipation, inflammations, fever, ascites,
bronchitis, cough, skin diseases, colic and lumbago. In Yunani medicine, castor root is used as
purgative and for skin diseases, the leaves are used to increase breastmilk production and for burns,
the seeds and the oil from them are purgative, useful in liver troubles, pains, lumbago, boils, piles,
ingrowworm, inflammations, ascites, asthma, rheumatism, dropsy and amenorrhea (reviewed in [9]).
Ground castor seeds or leaf paste were applied in veterinary medicine to heal sprains, swellings and
wounds [10].

Castor bean is used in folk medicine widespread throughout the world and has been reported:
(i) as a galactogogue on the Mediterranean coasts of Europe, where fresh leaves or leaf juice are
applied on the puerperal breast to promote lactation; (ii) as a remedy for various articular, cutaneous
or ocular diseases in Africa, where crushed seeds or oil, sometimes in combination with other plants,
are spread or rubbed on the sickling part of the body, or a root decoction is drunk to induce uterine
contraction as an abortive; (iii) as a medicament to cure erysipelas, flu, inflammation of the womb
and stomach aches in the Caribbean, where a leaf poultice is recommended; (iv) as an anthelmintic
or a purgative in Brazil where the seed oil is orally assumed or locally applied with the purpose of
contrast the hair loss or of healing wounds or burns (reviewed in [11]).

The laxative and abortifacient activities of castor seeds were attributed to the activation of
intestinal and uterine smooth-muscle cells via prostaglandin EP3 receptors induced by ricinoleic acid
[12]. Castor oil-induced diarrhea can be antagonized by hexane extract of Citrus limon peel that
activates antisecretory and antimotility mechanisms through the β adrenergic system [13]. The
purgative and anthelmintic actions of the oral ingestion of castor seeds, at least in part, could also be
ascribed to the irritative effect caused to intestine by ricin, as reported in toxicological studies
(reviewed in [14]). In addition, the antiflogistic action of castor bean could be related to the high
toxicity of ricin to macrophagic cells, which are responsible of producing inflammatory cytokines
(reviewed in [15]). This effect, together with the anti-pathogen activity of ricin, could promote healing
of the lesions, thus justifying its use in the treatment of various skin affections.

2. The ricin story

In the past centuries, castor oil had many uses; it was obtained by crushing the seeds and the
process produced a significant amount of residual press cake that was known to contain a highly
toxic component. Castor seed toxicity began to be investigated at the end of nineteenth century at
Schmiedeberg's laboratory in Strasbourg. The toxic component of Ricinus could be extracted with water and precipitated with alcohol, but it lost its toxic activity through heating, treatment with strong acid or repeated precipitation with alcohol. In 1887, Dixson supposed that the toxicity of Ricinus was due to a protein [16]. However, there were still many doubts whether the seed toxicity was due to a protein or a glycoside (reviewed in [17]). The problem was solved at the Medical Faculty of Dorpat (now Tartu) where an extremely toxic protein was partially purified from castor seed or press cake and named ricin in the doctoral thesis written by Hermann Stillmark under the supervision of Prof. Rudolf Kobert [18]. Stillmark noticed the agglutinating activity of ricin on red blood cells that was mistakenly believed to be the cause of ricin toxicity until the agglutinin was separated from the toxin [19].

Paul Ehrlich began his experiments in immunology by feeding mice with small amount of ricin or abrin, another similar plant toxin, until they were accustomed and became resistant to the toxin used, still remaining sensitive to the other toxin. The immunization was strictly specific, started after a few days and persisted at least for several months [20,21]. He was successful in the production of antisera against abrin and ricin and in the determination of antibody titer in serum and milk. Ehrlich drew animal experiments that clarified the transmission of passive immunity from mother to offspring through the transplacental transfer of antibodies and the breastfeeding. He investigated the dynamics of the antibody response and was the first to envisage the presence of binding sites on the cell surface (reviewed in [22]). These studies together with those on the immunity to bacterial toxins led him to formulate his side-chain theory of antibody formation and to win, in 1908, the Nobel Prize [23].

Interest in ricin was rekindled when the anticancer activity of this toxin on Ehrlich ascites cells in a mouse model was published [24]. A strong inhibition of protein synthesis by ricin was observed in cultures of both Ehrlich ascites tumour cells and Yoshida ascites hepatoma cells. The inhibition of protein synthesis by ricin requires more time in rat liver than in neoplastic cells [36]. The prospect of a possible use in cancer therapy induced to investigate which part of the proteosynthetic machinery was damaged and how the toxin managed to enter the cell to reach its target. In this paper, we highlight the milestones of research on ricin, with particular regards to its enzymatic activity, structure, cytotoxicity, toxicity for animals and humans and employ as immunotoxins, used in experimental models and in clinical trials. The main milestones are shown in Figure 1.

2.1. Ricin structure

The first information about the bi-chain nature of ricin structure dates to the early 70s, when it was shown that ricin was composed by two chains, A (active) and B (binding), linked together through a disulphide bond [26,27]. In the same period, the complete primary sequence of the ricin A and B chains was determined [28,29]. Ricin holotoxin structure was solved for the first time at 2.8 Å resolution (Figure 1) [30]. This pioneering work demonstrated that ricin A chain was a globular protein folded into three domains all contributing to the active site, while the B chain lectin folded into two domains, each binding lactose in a shallow cleft. The interface between the A and B chains showed some hydrophobic contacts in which proline and phenylalanine side chains played a prominent role. Four years later, the same researchers refined ricin structure at 2.5 Å (Figure 2a), allowing a more detailed molecular description of the holotoxin and of the separated A and B chains [31-33].
Figure 1. The main milestones of ricin research.

Ricin A chain was described as a globular protein consisting of 267 amino acids and organized in 8 α-helices and 8 β-strand structures. Ricin B chain consists of 262 amino acids and two homologues domains, each containing a lactose binding site and several areas of amino acid homology, possibly derived from a gene duplication. In 1995, after purification of a complex of ricin A chain cross-linked to the ribosome, it was found the binding of ricin A chain with the ribosomal proteins L9 and L10e [34,35].

The knowledge of tridimensional structure of ricin yielded more information on its active site. Studies based on the formation of complexes between the A chain, both native and recombinant, and adenine-containing nucleotides allowed the identification of key residues in enzymatic activity. In particular, Tyr80, Tyr123, Glu177, Arg180 and Trp211 were found to form the binding site for adenine (Figure 2b) [30,36]. In the 90s, the molecular mechanism of de-adenilation was hypothesized: adenine is sandwiched between Tyr80 and Tyr123 in a π stacking interaction; the N3 of adenine is protonated by Arg180, promoting the C1’-N9 bond breaking thus forming an oxycarbene moiety on the ribose (Figure 2c) [36,37]. This transition state is stabilized by Glu177; a water molecule lies on the opposite side of the sugar ring from adenosine, which will be polarized by Arg180 to a hydroxide character that rapidly attacks the sugar carbon completing the reaction.
2.2. Ricin enzymatic activity

The introduction of a cell-free system utilizing a lysate from rabbit reticulocytes [38] helped to clarify that ricin inhibited the peptide chain elongation (Figure 1) [27]. The two polypeptides showed different properties: the A chain possessed the toxic activity, while the B chain was a galactose-specific lectin binding the cell surface [26]. Treating the toxin with reducing agents, it resulted more active in inhibiting cell-free protein synthesis [39]. Firstly, the target of the toxic action was identified as the ribosome (Figure 1), then as the 60 S subunit of eukaryotic ribosome [40], which became unreactive toward elongation factors [41]. The toxin was found to interfere with the interaction of elongation factors with the ribosomes and their elongation-factor-dependent GTPase activity [41,42]. The A-chain molecule resulted very active on its substrate and it was calculated that one molecule can inactivate 2000 ribosomes/min, with a $K_m$ of 0.1-0.2 mM [43].

In addition to ricin, several other plant proteins have been identified to possess a similar protein synthesis inhibiting action. Most of them had a single polypeptide chain similar to the A chain of ricin. They were called Ribosome-Inactivating Proteins (RIPs) (reviewed in [44,45]).

The already supposed enzymatic nature of ricin A chain was finally demonstrated in 1987 by Endo and Co-workers, which discovered that ricin A-chain cleaved the N-glycosidic bond of an adenine residue, A4324 in rat 28 S RNA, from the ribose of a highly conserved ribosomal RNA single-stranded loop involved in the binding of elongation factors (Figure 1). The toxin did not directly break the RNA chain, but the depurinated RNA was susceptible to hydrolysis [46,47]. Consequently, ricin activity was identified as an rRNA N-glycosidase [EC 3.2.2.22].
After, it was demonstrated that the enzymatic activity of RIPs was broader than previously described. All tested RIPs were able to deadenylate DNA, in addition to rRNA, and some of them were also able to deadenylate different other polynucleotide substrates, releasing adenine from the sugar phosphate backbone of polynucleotide substrates (Figure 1) [48,49]. For this reason, the name of adenine polynucleotide glycosylase was proposed for RIPs. Thus, the ability of acting on various substrates and extensively depurinating some of them suggested that the protein synthesis inhibition could be only one of the ways of RIP-mediated cell killing. Ricin resulted able to deadenylate rRNA, DNA (chromatin and naked) and also poly(ADP-ribosyl)ated poly(ADP-ribose) polymerase, an enzyme involved in DNA repair [48,50]. Furthermore, it was observed that many RIPs were able to cleave more than one adenine; although ricin was able to detach few adenines from the DNA (tens), less than some single-chain RIPs (thousands). The hypothesis that ricin could act directly on DNA in cellular models was strengthened by the evidence that damage to nuclear DNA, consistent with the enzymatic activity (adenine release) on DNA in cell-free systems, was concomitant with protein synthesis inhibition and preceded apoptosis [51].

2.3. Ricin cellular uptake, routing and toxicity

Starting from the mid-70s, several research groups focused on ricin binding and internalization studies, demonstrating that the interaction of ricin with the cell started from the binding of the B chain to galactosyl residues on cell surface, allowing the access to endosomal compartment [52]. Ricin binds to both glycolipids and glycoproteins with terminal galactose. Since ricin binds to a variety of different molecules, it seems to be internalized by different endocytic pathways as well as to use different pathways to reach the Golgi apparatus and to intoxicate the cell. In HeLa cells, about 107 binding sites were found for ricin, but only small amount of the bound toxin reached the Golgi network and participated to the cell intoxication [52].

Firstly, it was reported that ricin entered into cytoplasm through clathrin-dependent endocytosis [53]. Afterwards, it became clear that also clathrin-independent mechanisms were involved [54]. After cell uptake, ricin is delivered to early endosomes, from where most of protein molecules are recycled back to the cell surface or delivered, via late endosomes, to lysosomes for proteolytical degradation. A small amount of not degraded ricin is addressed within the trans-Golgi network [55]. The involvement of Golgi complex in ricin routing was confirmed using different Golgi-disrupting agents, such as brefeldin A, monensin, etc. In fact, the pretreatment with these agents inhibited the cytotoxic effects of ricin [56]. It was demonstrated that ricin was cycled from Golgi to endoplasmic reticulum via coatomer protein 1 (COP-1)-coated vesicles [57], although it was later proved that also the COP-1-independent pathway could be involved [58].

The complete elucidation of intracellular ricin traffic occurred when it was demonstrated that, after reaching the endoplasmic reticulum, the two ricin chains were separated, and the A chain was retro-translocated through the quality control pathway delivering misfolded proteins to cytosol (Figure 1) [59]. Recently it has been demonstrated that cholesterol rafts are required for Golgi transport of ricin; instead, glycosphingolipids seem to be not required (reviewed in [60]).

The portion of A chain that quickly refolded, thus avoiding ubiquitination and proteosomal degradation, was able to reach its intracellular target (reviewed in [61]). It was estimated that one molecule of active ricin that arrives to its substrate is enough to kill one cell [62].

The discovery that ricin and some related toxins may be retrogradely transported along neuronal processes (Figure 1) [63] opened a new field of research in neurobiology and this property has been exploited for the selective destruction of neuron bodies.

Different cell types showed variable levels of sensitivity to ricin (reviewed in [14]), possibly because of the mannose receptor expression on cell surface and the endocytosis efficacy. Ricin results one of the most toxic plant toxins on cell lines with IC50s (concentration inhibiting protein synthesis by 50%) ranging from less than 0.1 to 1 pM [26,64-66]. However, it must be taken into account that it
is very difficult to make a direct comparison of the data available in literature about ricin cytotoxicity, because of the differences in the experimental approaches and technical conditions.

The polynucleotide depurinating activity of RIPs suggests the possibility of a wider toxic action on many biological substrates, not excluding the induction of oxidative stress, all together justifying the existence of more than one death mechanism induced by ricin and other RIPs, such as apoptosis and necroptosis (Figure 1) [64,67].

3. Ricin toxicity in humans and animals

On one hand, ricin has been studied for bio-medical applications, exploiting the ability of the A-chain to kill target cells once linked to a monoclonal antibody, as below described in the immunotoxins chapter. On the other hand, ricin has attracted nefarious interests, with a history of military, criminal and terrorist uses [68].

The acute toxicity of ricin is highly variable depending on the animal species and the strain. The pathological effects and subsequent clinical signs of ricin intoxication depend also on the route of exposure, as this dictates the subsequent tissue distribution of the toxin. Following intravenous or intramuscular administration, lesions eventually develop in the spleen, liver and kidney whilst the lung remains unaffected. After oral ingestion, the gastrointestinal tract is severely affected. Inhalational exposure produces effects that are mainly confined to the respiratory tract [69].

Main data on animal toxicity derived from laboratory experiments in rodents, principally rat and mouse. Oral administration of ricin was reported to give Lethal Dose for 50% of animals (LD₅₀) of 20-30 mg/kg in rat and 15–35 mg/kg in mouse [70-72]. For intravenous, inhalation and intraperitoneal routes, toxicity is approximately 1000-fold higher than for oral route, with LD₅₀ values in mouse of 2-10 μg/kg, 3-5 μg/kg and 22 μg/kg, respectively [70,73]. The lower toxicity of ricin after oral exposure is due to the protein destruction in the lumen of the intestinal tract [74,75]. Ricin acts in a time- and concentration-dependent manner. Notably, there is a time delay of about 10 h before death occurs even when very high doses are applied [76].

Oral toxicity. In human, most intoxications occurred accidentally or voluntarily with the ingestion of castor seeds; only few cases of intentional absorption of castor bean extracts have been documented in suicide attempts [76]. Whole-ingested beans can pass intact through the gastrointestinal tract, whereas chewing facilitates ricin release. Also, it has been reported that the seed can act as “timed-release” capsule for the toxin, allowing its release in the lower bowel, where it causes more damage [72]. After ingestion, vomiting, diarrhea and abdominal pain are common symptoms. Massive gastrointestinal fluid and electrolyte loss are described, often complicated by hematemesis or melaena. Finally, hypovolemic shock and multiorgan failure occur, which particularly involves spleen, liver and kidney [77,78].

Despite the high number of intoxicated subjects with castor beans, it is quite difficult to calculate LD values for ricin in humans. In fact, the effective ingested ricin dose can only be supposed, because of ricin content variations depending on size, weight and moisture of seeds, as well as on cultivar, region, season and plant growth stage. Moreover, in intoxicated subjects, it must be taken into account the degree of mastication, stomach content, age and comorbidities, parameters that are obviously more heterogeneous compared to experimental poisoning of animals. Considering all these parameters, the fatal oral dose of ricin in humans has been estimated to range from 1 to 20 mg/kg (approx. 5-10 beans) [70,79].

Inhalation toxicity. No data are available for human ricin uptake by inhalation. In non-human primates, LD₅₀ has been estimated 5–15 μg/kg depending on aerosol particle size. Inhalation of particles that are able to penetrate deeply into the lungs (1-5 μm diameter) displays much more toxicity than larger particles [72,80]. Inhalation of ricin causes slow onset of respiratory distress (difficulty breathing), coughing, fever, pulmonary lesions and edema. Intoxicated animals develop fibrinopurulent necrotizing pneumonia accompanied by necrotizing lymphadenitis, typically after a dose-dependent delay of 8-24 hours. Death occurs for respiratory failure due to massive alveolar fluid...
accumulation. Liver, kidney and small intestines appear congested although little histologic changes are shown [72,80,81].

Parenteral toxicity. Data regarding parenteral ricin intoxication derive mainly from animal lab. By injection, mice had an LD₅₀ of 3-5 μg/kg by intravenous and 22 μg/kg by subcutaneous route [82], rabbits had LD₅₀ 0.5 μg/kg by intravenous and 0.1 μg/kg by intramuscular route, while guinea pigs had LD₅₀ <1.1 μg/kg by intravenous and 0.8 μg/kg by intramuscular route [83]. Human data derive from few cases of suicide or murder, or their attempt; the most known episode is the assassination of the Bulgarian dissident Georgi Markov who in 1978 died 3 days after being stabbed probably with an umbrella loaded with a ricin-containing pellet (Figure 1) [84].

By parenteral administration, immediate local pain at the injection site is reported, followed by general weakness within 5 hours. The following symptoms, that are general and maybe similar to sepsis (fever, headache, dizziness, anorexia, nausea, vomiting, hypotension, abdominal pain), can be delayed for as much as 10 to 12 hours, even with high doses. Usually local tissue damage at the site of the injection was observed. Laboratory abnormalities included elevated liver transaminases, amylase and creatinine kinase, hyperbilirubinemia, myoglobinuria and renal insufficiency. The clinical course may progress to multisystem organ failure. Preterminal complications included gastrointestinal hemorrhage, hypovolemic shock and renal failure [78,84].

4. Bioterrorism and environmental toxicity

Ricin is currently monitored as a Schedule 1A of the Chemical Weapons Convention (CWC) and is a Category B substance under the Biological and Toxins Weapons Convention (BTWC) [80]. Despite its toxicity, ricin is less potent than other agents such as botulinum neurotoxin or anthrax. It has been estimated that eight tons of ricin would have to be aerosolized over a 100 km² area to achieve about 50% casualty, whereas only kilogram quantities of anthrax spores would cause the same effect [85]. Thus, deploying an agent such as ricin over a wide area, although possible, becomes impractical from a logistics standpoint. However, the availability of castor beans and the quite simple procedure for rough ricin purification have attracted criminal and terrorist interest for small scale biocrimes or to cause collective media-driven alarm [80].

From castor seeds a nontoxic oil can be extracted that find a multitude of uses in many sectors, including cosmetic, pharmaceutic, mechanical and chemical industry. The castor oil production is increasing worldwide because of its versatile application, low cost, availability and biodegradability. In addition, the oil-free seed pulp can be used in agriculture as a natural fertilizer [86], although the processing of castor seeds requires great caution due to the high allergenicity [87,88] and extreme toxicity [76] of their protein fraction, represented above all by ricin. World production of castor oil increased from 0.8 million tons in 2000 [89] to 1.21 million tons in 2014 [90], with a castor seed production of 1.49 million tons in 2017 [91]. Leading producing countries are India, with over 80% of the global yield, Mozambique, China, Brazil, Myanmar, Ethiopia, Paraguay and Vietnam [92]. The oil makes up about 50% of the weight of the seeds and is mostly constituted of ricinoleic acid (90%), with minor amounts of dihydroxystearic, linoleic, oleic and stearic acids. Ricin isoforms and the alkaloid ricinine, are not transferred to the oil fraction during extraction, which can be performed by cold or warm pressing, but remain in the seed cake [93,94].

Castor bean meal press cake or other residues of the castor oil production have been employed as a protein source for feed or fertilizer, but their use is very limited by the ricin toxicity [76]. In 2008, the European Food Safety Agency defined ricin as undesirable substance in animal feed. Ricinus derived material should be appropriately inactivated through physical and/or chemical methods to guarantee animal and human health [95]. Nevertheless, many accidental poisonings are still reported for animals eating improperly detoxified fertilizer or other agricultural products containing castor derived material [76,94].
In order to block the toxic action of ricin, different strategies have been evaluated: vaccines, inhibitors and passive immunity. Vaccines against ricin with the consequent production of neutralizing antibodies did not give satisfactory results in vivo (reviewed in [96]). Inhibitors of ricin can act blocking the active site or as substrate analogue; also in this case, the available data are limited to in vivo experiments [97]. More recently, inhibitors of cell routing have been developed, sometimes giving promising results also in vivo [98,99].

To date passive immunity has been proven the only effective strategy for treating intoxications caused by ricin. The delay in the appearance of signs of intoxication makes confirmation of exposure, diagnosis of intoxication and the subsequent medical response technically and logistically challenging. The development of anti-ricin sera or antibodies, effective even when used several hours after toxin exposure, represents a step forward in treatment of ricin intoxication, as it increases the time window of intervention (WOO, window of opportunity). Many authors described effective post-exposure treatment of ricin intoxication with specific antibodies, but with a limited WOO (~8 h) [100-103]. Other authors reported a survival between 50 and 89% of mice treated with anti-sera 24 hours after intoxication [104-106]. Once internalized into the cells, ricin cannot be neutralized by antibodies, thus limiting the therapeutic window. However, Whitfield et al. in 2017 reported 100% protection in aerosolized ricin-treated mice with a single administration of a F(ab’)2 polyclonal ovine antitoxin given 24 h post-intoxication [107]. Even when performed in the same animal species, comparison between diverse experiments is often difficult, due to the different toxin dose and route of administration utilized. Moreover, there are few data about correlation between the antitoxin dose required for protection and the WOO.

5. Ricin-containing immunotoxins

Many researchers tried to exploit the high ricin cytotoxicity for medical purposes to eliminate pathological cells. Although ricin possesses highly efficient cell killing mechanisms, it lacks selectivity towards cell targets. In order to increase selectivity, it has been explored the possibility of linking ricin to carriers specific for targets on unwanted cells. The most widely used carriers are antibodies and the corresponding conjugates are referred to as immunotoxins (ITs).

The first IT, created in 1976 by Moolten and co-workers, was made by Ricin Toxin-A chain (RTA) linked to a rat tumor-specific antibody against a rat lymphoma, namely (C58NT)D (Figure 1) [108]. So far, a multitude of preclinical and clinical studies have shown the potential use of several ricin-ITs towards different cancer types, from hematological to solid ones, and towards normal cells, unwanted because responsible of a pathological state (reviewed in [109,110]). Different approaches have been used, over time, to generate ITs. In the first strategy, ITs were composed by the antibody chemically linked to the entire RIP that were used for in vitro and in vivo studies showing high cytotoxicity [111]. Despite the high in vitro efficiency, the relevant non-specific toxicity reported in vivo, due to the characteristics of the lectin chain, brought researchers to sterically block, chemically modify or remove the B chain, thus balancing toxicity and specificity. In 1985, Weil-Hillman and colleagues tested an anti-Mr 67,000 protein linked to either blocked-chain B ricin or RTA, in a nude mouse model reporting interesting results in vitro but not in vivo [112]. The 80s were years of great ferment for molecular biology and genetic engineering, paving the way for the second generation of ITs. Many researchers tried to improve the IT penetration in tumor mass by reducing the antibody size, using antigen-binding (Fab) or variable (Fv) fragments instead of entire antibodies. In 1988, Ghetti and colleagues created a new IT composed by Fab’ fragments conjugated to chemically deglycosylated RTA (dgA) [113]. Few years later, they used an anti-CD122-dgA IT in SCID-Daudi mice, showing promising results since the IT was able to specifically kill tumor cells in vivo, extending the mean survival time up to 57.9 +/- 3.8 days [114]. Moreover, FitzGerald and co-workers described the antitumor activity of recombinant RTA (rRTA) linked to anti-mouse transferrin receptor in a nude mouse model of human ovarian cancer. Animals treated with IT extended life span...
from 45 (lower doses) to 70/80 days (higher doses) [115]. Finally, in 1997 the first ricin-containing recombinant immunotoxin (rIT) was obtained through the expression of a fusion gene composed by sequences encoding anti-CD19-FVS191 (single-chain Fv), cathepsin D proteinase digestion site and rRTA. In this work, authors compared the cytotoxicity of the rIT with the chemical linked IT, evidencing that only the latter was toxic in target cells [116]. About 20 ricin-ITs have been tested in Phase I, II and III clinical trials to treat patients with either hematological or solid tumors, transplant rejection and GvHD. One of the first Phase I clinical trials has been conducted by Spitzer and colleagues in 1987 (Figure 1), in which they obtained promising results by treating 22 metastatic malignant melanoma patients with an IT composed by murine monoclonal anti-melanoma antibody coupled to RTA (XOMAZYME-MEL) [117-119]. Additionally, ITs were also exploited for the treatment of autoimmune disease. Indeed, anti-CD5/RTA was the first IT to be used in clinical trials for therapy of autoimmune diseases, such as rheumatoid arthritis, systemic lupus erythematosus and insulin-dependent diabetes mellitus [120,121].

A different approach can be represented by the construction of nanoparticles, in which ricin is genetically fused to carrier peptides that are able not only to recognize specific cellular target, but also to auto assemble, as stable nanoparticles, thus increasing the toxin-concentration into the targeted site [110,122,123].

6. Conclusions

In conclusion, ricin is a highly cytotoxic plant protein and has been of great utility to develop a number of anti-cancer immunotoxins. The ability of ricin, and of some other RIPs, to act on multiple molecular target inside the cell, thus triggering different death pathways, makes it more attractive for cancer treatment than conventional chemotherapy, in which one of the major problems is the rise of resistant cells [67,124]. However, ricin-containing ITs also exhibited many limitations like unspecific toxicity, organ toxicity (mainly liver, kidney and vasculature), immunogenicity, fast removal from blood stream, and lysosomal degradation inside cells. As a result, despite of the significant efforts made over the past few years, ricin as therapeutic agent has not been achieved much impact at the clinical level. The challenge is still open, and the frontline research is directed towards “recombinant immunotoxins” and nanocarriers, or, probably, by other novel techniques represented by expressing active plant toxin genes in vector specific for tumor cells [110,125].

Although ricin is not enough toxic to hypothesize a use over a wide area for terrorist purposes, the availability of castor beans and the quite simple procedure for rough ricin purification have stimulated the interest of criminals and terrorists for small-scale biocrimes. This justifies the researchers’ efforts to obtain always more fast and sensitive ricin detection tests. In addition, the study of inhibiting or neutralizing molecules and the timing of clinical events in case of ricin intoxication could lead to the definition of one or more validated therapies.

Finally, the use of castor bean derivatives should be carefully monitored because of the potential presence of active ricin. In fact, the large use of these products in agriculture, without an effective ricin inactivation, caused several cases of animal intoxication and can be hazardous for human health.

Author Contributions: All the authors contribute to collect the literature, write and revise the paper.

Acknowledgments: This work was supported by funds for selected research topics from the Alma Mater Studiorum—University of Bologna and by the Pallotti Legacies for Cancer Research.

Conflicts of Interest: The authors declare no conflict of interest.

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