Nuclear Fusion Catalyzed by Doubly Charged Scalars:
Implications for Energy Production

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A number of popular extensions of the Standard Model (SM) of particle physics predict the existence of doubly charged scalar particles $X^{\pm \pm}$. Such particles may be long-lived or even stable. If exist, $X^{--}$ could form atomic bound states with light nuclei and catalyze their fusion by essentially eliminating the Coulomb barrier between them. Such an $X$-catalyzed fusion (XCF) process does not require high temperatures or pressure and may have important applications for energy production. A similar process of muon-catalyzed fusion ($\mu$CF) has been shown not to be a viable source of energy because of the sticking of negative muons to helium nuclei produced in the fusion of hydrogen isotopes, which stops the catalytic process. We analyze XCF in deuterium environments and show that the X-particles can only stick to $^6$Li nuclei, which are produced in the third-stage reactions downstream in the catalytic cycle. The corresponding sticking probability is very low, and, before getting bound to $^6$Li, each X-particle can catalyze $\sim 3.5 \cdot 10^9$ fusion cycles, producing $\sim 7 \cdot 10^4$ TeV of energy. We also discuss the ways of reactivating the X-particles from the Coulomb-bound ($^6$Li$X$) states, which would allow reusing them in XCF reactions.

I. INTRODUCTION

A number of popular extensions of the SM of particle physics predict the existence of doubly charged scalar particles $X^{\pm \pm}$. These include the Type-II seesaw [1–6] and the Zee-Babu [7, 8] models of neutrino masses, the Left–Right model [9–11], the Georgi–Machacek model [12–16], the 3-3-1 model [17, 18] and the little Higgs model [19]). Doubly charged scalars appear also in simplified models, in which one merely adds such scalars in a gauge invariant way in various representations of the SM gauge group $SU(2)_L$ to the particle content of the SM. The Lagrangian of the model is then complemented by gauge-invariant interaction terms involving these new fields [20, 21].

Doubly charged scalars may be long-lived or even stable [21–24]. As the simplest example, one can add to the SM an uncolored $SU(2)_L$-singlet scalar field $X$ with hypercharge $Y = 2$ [21]. The corresponding doubly charged particle will couple to the neutral gauge bosons $\gamma$ and $Z^0$ and may also interact with the SM Higgs boson $H$ through the $(H^\dagger H)(X^\dagger X)$ term in the Higgs potential. Gauge invariance allows, in addition, the Yukawa coupling of $X$ to right-handed charged leptons, $h_X l_R l_R X + h.c.$ This is the only coupling that makes the $X$-particles unstable in this model; they will be long-lived if the Yukawa coupling constants $h_X$ are small. The Yukawa coupling of $X$ may be forbidden by e.g. $Z_3$ symmetry $X \rightarrow -X$, in which case the $X$-scalars will be stable. Doubly charged scalar particles are being actively searched for experimentally, but up to now have not been discovered. For discussions of current experimental constraints on the doubly charged particles and of the sensitivities to them of future experiments see [21–27] and references therein.

In addition to interesting particle-physics phenomenology, doubly charged scalars may have important implications for cosmology. In this paper we will, however, consider another aspect of their possible existence. As we shall demonstrate, doubly charged particles can catalyze fusion of light nuclei, with potentially important applications for energy production. The negatively charged $X^{--}$ (which we will hereafter simply refer to as $X$) can form atomic bound systems with the nuclei of light elements, such as deuterium, tritium or helium. One example is the antihelium-like ($ddX$) atom with the $X$-particle as the “nucleus” and two deuterons in the 1s atomic state instead of two positrons. (Here and below we use the brackets to denote states bound by the Coulomb force).

As $X$ is expected to be very heavy, the size of such an atomic system will in fact be determined by the deuteron mass $m_d$ and will be of the order of the Bohr radius of the $(dX)$ ion, $a_d \simeq 7.2$ fm. Similar small-size atomic systems ($NN^*X$) can exist for other light nuclei $N$ and $N'$ with charges $Z \leq 2$.

Atomic binding of two nuclei to an $X$-particle brings them so close together that this essentially eliminates the necessity for them to overcome the Coulomb barrier in order to undergo fusion. The exothermic fusion reactions can then occur unhindered and do not require high temperature or pressure. The $X$-particle is not consumed in this process and can then facilitate further nuclear fusion reactions, acting thus as a catalyst.

This $X$-catalyzed fusion mechanism is to some extent similar to muon catalyzed fusion ($\mu$CF) [28–35], see [36–42] for reviews), in which the role of the catalyst is played by singly negatively charged muons. $\mu$CF of hydrogen isotopes was once considered a prospective candidate for cold fusion. However, already rather early in its studies it became clear that $\mu$CF suffers from a serious shortcoming that may prevent it from being a viable mechanism of...
energy production. In the fusion processes, isotopes of helium are produced, and there is a chance that they will capture on their atomic orbits the negative muons present in the final state of the fusion reactions. Once this happens, muonic ions (3Heµ) or (4Heµ) are formed, which, being positively charged, cannot catalyze further fusion reactions. This effect is cumulative; the sticking to helium nuclei thus eventually knocks the muons out from the catalytic process, i.e. the catalytic poisoning occurs.

Out of all μCF reactions, the d−t fusion has the smallest muon sticking probability, ωs ∼ 10−2. This means that a single muon will catalyze ~100 fusion reactions before it gets removed from the catalytic process. The corresponding total produced energy is ∼1.7 GeV, which is at least a factor of five smaller than the energy needed to produce and handle one muon [32]. In addition, muon’s short lifetime makes it impractical to try to destroy the produced (3Heµ) or (4Heµ) bound states by irradiating them with particle beams in order to reuse the released muons. These considerations have essentially killed the idea of using μCF for energy production.

There were discussions in the literature of the possibility of energy generation through the catalysis of nuclear fusion by hypothetic heavy long-lived or stable singly charged [34, 43-45] or fractionally charged [46] particles. However, it has been shown in [34, 44, 45] that these processes suffer from the same problem of catalytic poisoning as μCF, and therefore they cannot be useful sources of energy. In particular, in ref. [44] it was demonstrated that reactivation of the catalyst particles by irradiating their atomic bound states with helium nuclei by neutron beams, as suggested in [46], would require beams that are about nine orders of magnitude higher than those currently produced by most powerful nuclear reactors.

In this paper we consider the fusion of light nuclei catalyzed by doubly negatively charged X-particles and demonstrate that, unlike μCF, this process may be a viable source of energy. We analyze X-catalyzed fusion (XCF) in deuterium environments and show that the catalytic poisoning may only occur in this case due to the sticking of X-particles to 6Li nuclei, which are produced in the fusion reactions of the third stage. The corresponding sticking probability is shown to be very low, and, before getting bound to 6Li, each X-particle can catalyze ∼ 3.5 · 109 fusion cycles, producing ∼ 7 · 104 TeV of energy. To the best of the present author’s knowledge, nuclear fusion catalyzed by doubly charged particles has never been considered before.

II. X-CATALYZED FUSION IN DEUTERIUM

We will be assuming that X-particles interact only electromagnetically, which in any case should be a very good approximation at low energies relevant to nuclear fusion. Let X-particles be injected in pressurized D2 gas or liquid deuterium. Being very heavy and negatively charged, the X-particles can easily penetrate D2 molecules and D atoms, dissociating the former and ionizing the latter and losing energy on the way. Once the velocity of an X-particle becomes comparable to atomic velocities (v ∼ 2e2/h ∼ 10−2 c), it captures a deuteron on a highly excited atomic level of the (dX) system, which then very quickly de-excites to its ground state, mostly through electric dipole radiation and inelastic scattering on the neighboring deuterium atoms. As the (dX) ion is negatively charged, it swiftly picks up another deuteron to form the (ddX) atom. The characteristic time of this atomic phase of the XCF process is dominated by the X moderation time and is ∼ 10−16 s at liquid hydrogen density N0 = 4.25 × 1022 nuclei/cm3 and T ∼ 20 K and about 10−7 s in deuterium gas at 0°C and pressure of one bar (see Appendix A).

After the (ddX) atom has been formed, the deuterons undergo nuclear fusion through several channels, see below. Simple estimates show that the fusion rates are many orders of magnitudes faster than the rates of the atomic formation processes. That is, once (ddX) [or similar (NN’X)] atoms are formed, the fusion occurs practically instantaneously. The time scale of XCF is therefore determined by the atomic formation times. The rates of the fusion reactions, however, determine the branching ratios of various fusion channels, which are important for the kinetics of the catalytic cycle.

At the first stage of XCF in deuterium two deuterons fuse to produce 3He, 3H or 4He. In each case there is at least one channel in which the final-state X forms an atomic bound state with one of the produced nuclei. Stage I fusion reactions are

(1a) (ddX) → 3He + n + X (Q = 2.98 MeV, 29.1%)  
(1b) (ddX) → (3HeX) + n (Q = 3.89 MeV, 19.4%)

(2a) (ddX) → 3H + p + X (Q = 3.74 MeV, 34.4%)  
(2b) (ddX) → (3HX) + p (Q = 4.01 MeV, 6.2%)

(2c) (ddX) → 3H + (pX) (Q = 3.84 MeV, 0.5%)

(3a) (ddX) → 4He + γ + X (Q = 23.6 MeV, 4·10−9)  
(3b) (ddX) → (4HeX) + γ (Q = 24.7 MeV, 3·10−8)

(3c) (ddX) → 4He + X (Q = 23.6 MeV, 10.4%)

Here in the parentheses the Q-values and the branching ratios of the reactions are shown. In evaluating the Q-values we have taken into account that the atomic binding of the two deuterons to X in the initial state reduces Q, whereas the binding to X of one of the final-state nuclei increases it. As the Bohr radii of most of the X-atomic states we consider are either comparable to or smaller than the nuclear radii, in calculating the Coulomb binding energies one has to allow for the finite nuclear sizes. We do that by making use of a variational approach, as described in Appendix A 2.

The rates of reactions (1b), (2b), (2c) and (3b) with bound X-particles in the final states are proportional to the corresponding X-particle sticking probabilities,
\[ \omega_\alpha. \] The existence of such channels obviously affects the branching ratios of the analogous reactions with free \( X \) in the final states.

Radiative reactions (3a) and (3b) have tiny branching ratios, which is related to their electromagnetic nature and to the fact that for their \( X \)-less version, \( d + d \to ^4\text{He} + \gamma \), transitions of E1 type are strictly forbidden. This comes about because the two fusing nuclei are identical, which, in particular, means that they have the same charge-to-mass ratio. This reaction therefore proceeds mainly through E2 transitions \([35]\). When the deuterons are bound to \( X \), the strict prohibition of E1 transitions is lifted due to possible transitions through intermediate excited atomic states.\(^1\) However, as shown in Appendix A 3b, the resulting E1 transitions are in this case heavily hindered and their rates actually fall below the rates of the E2 transitions.

Reaction (3c) is an internal conversion process. Note that, unlike for reactions (1a) - (3b), the \( X \)-less version of (3c) does not exist: the process \( d + d \to ^4\text{He} \) is forbidden by kinematics. For the details of the calculation of the rate of reaction (3c) as well as of the rates of the other discussed in this paper reactions, see Appendix B. The relevant \( Q \)-values of the reactions and sticking probabilities are evaluated in Appendices A 2 and A 3, respectively.

The final states of reactions (1a), (2a), (3a) and (3c) contain free \( X \)-particles which are practically at rest and can immediately capture deuterons of the medium, forming again the \((ddX)\) atoms. Thus, they can again catalyze \( d-d \) fusion through stage I reactions (1a)-(3c). The same is also true for the \( X \)-particles in the final state of reaction (2c) which emerge being bound to protons. Collisions of \((pX)\) with deuterons of the medium lead to fast replacement of the protons by deuterons through the exothermic charge exchange reaction \((pX)+d \to (dX)+p\) with the energy release \(\sim 90\ \text{keV}\) (see Appendix A 1b). The produced \((dX)\) ion then picks up a deuteron to form the \((ddX)\) atom, which can again participate in stage I reactions (1a)-(3c).

The situation is different for the \( X \)-particles in the final states of reactions (1b) and (2b) forming the bound states with \(^3\text{He}\) and \(^3\text{H}\), respectively. They can no longer directly participate in stage I \( d-d \) fusion reactions. However, they are not lost for the fusion process: the produced \((^3\text{He}X)\) and \((^3\text{HX})\) can still pick up deuterons of the medium to form the atomic bound states \((^3\text{He}dX)\) and \((^3\text{HdX})\), which can give rise to stage II fusion reactions, which we will consider next.

Before we proceed, a comment is in order. While \((^3\text{HX})\) is a singly negatively charged ion which can obviously pick up a positively charged deuteron to form an \((^3\text{HdX})\) atom, \((^3\text{He}X)\) is a neutral \( X \)-atom. It is not immediately obvious whether it can form a stable bound state with \( d \), which, if exists, would be a positive ion. In the case of the usual atomic systems, analogous (though negatively charged) states do exist – a well-known example is the negative ion of hydrogen \(^-\text{H}^+\). However, the stability of \((^3\text{He}dX)\) cannot be directly deduced from the stability of \(^-\text{H}^+\); in the latter case the two particles orbiting the nucleus are identical electrons, whereas for \((^3\text{He}dX)\) these are different entities – nuclei with differing masses and charges. Nevertheless, from the results of a general analysis of three-body Coulomb systems carried out in \([47–49]\) it follows that the state \((^3\text{He}dX)\) (as well as the bound state \((^3\text{HdX})\) which we will discuss later on) should exist and be stable. For additional information see Appendix A 1c.

Once \((^3\text{He}X)\) and \((^3\text{HX})\), produced in reactions (1b) and (2b), have picked up deuterons from the medium and formed the atomic bound states \((^3\text{He}dX)\) and \((^3\text{HdX})\), the following stage II fusion reactions occur:

\[ (^3\text{He}dX) \to ^4\text{He} + p + X \quad (Q = 17.4\ \text{MeV}, 94\%) \quad (4a) \]
\[ (^3\text{He}dX) \to (^4\text{He}X) + p \quad (Q = 18.6\ \text{MeV}, 6\%) \quad (4b) \]
\[ (^3\text{HdX}) \to ^4\text{He} + (pX) \quad (Q = 17.5\ \text{MeV}, 3 \times 10^{-4}) \quad (4c) \]
\[ (^3\text{HdX}) \to ^4\text{He} + n + X \quad (Q = 17.3\ \text{MeV}, 96\%) \quad (5a) \]
\[ (^3\text{HdX}) \to (^4\text{He}X) + n \quad (Q = 18.4\ \text{MeV}, 4\%) \quad (5b) \]

In these reactions vast majority of \( X \) bound to \(^3\text{He}\) and \(^3\text{H}\) are liberated; the freed \( X \)-particles can again form \((ddX)\) states and catalyze stage I fusion reactions (1a)-(3c). The same applies to the final-state \( X \)-particles bound to protons, as was discussed above. The remaining relatively small fraction of \( X \)-particles come out of stage II reactions in the form of \((^3\text{He}X)\) atoms. Together with a very small amount of \((^4\text{He}X)\) produced in reaction (3b), they pick up deuterons from the medium and form \((^3\text{He}dX)\) states, which undergo stage III XCF reactions:

\[ (^4\text{He}dX) \to ^6\text{Li} + \gamma + X (Q = 0.32\ \text{MeV}, 10^{-13}) \quad (6a) \]
\[ (^4\text{He}dX) \to (^6\text{Li}X) + \gamma \quad (Q = 2.4\ \text{MeV}, 2 \times 10^{-8}) \quad (6b) \]
\[ (^4\text{He}dX) \to ^6\text{Li} + X \quad (Q = 0.32\ \text{MeV}, \approx 100\%) \quad (6c) \]

In these reactions, almost all previously bound \( X \)-particles are liberated and are free to catalyze again nuclear fusion through XCF reactions of stages I and II. The remaining tiny fraction of \( X \)-particles end up being bound to the produced \(^6\text{Li}\) nuclei through reaction (6b). However, as small as it is, this fraction is very important for the kinetics of XCF. The bound states \((^6\text{Li}X)\) are ions of charge +1; they cannot form bound state with positively charged nuclei and participate in further XCF reaction. That is, with their formation catalytic poisoning occurs and the catalytic process stops.

From the branching ratios of stage I, II, and III XCF reactions one finds that the fraction of the initially injected \( X \)-particles which end up in the \((^6\text{Li}X)\) bound state is \(\sim 2.8 \times 10^{-10}\). This means that each initial \( X \)-particle, before getting stuck to a \(^6\text{Li}\) nucleus, can catalyze \(\sim 3.5 \times 10^9\) fusion cycles.

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\(^1\) The author is grateful to M. Pospelov for raising this issue and suggesting an example of a route through which E1 transitions could proceed in reaction (3b).
Direct inspection shows that, independently of which sub-channels were involved, the net effect of stage I, II and III XCF reactions is the conversion of four deuterons to a $^6\text{Li}$ nucleus, a proton and a neutron:

$$4d \rightarrow ^6\text{Li} + p + n + 23.1 \text{ MeV}.$$ (7)

Therefore, each initial $X$-particle will produce about $7 \times 10^4$ TeV of energy before it gets knocked out of the catalytic process. It should be stressed that this assumes that the $X$-particles are sufficiently long-lived to survive during $3.5 \times 10^9$ fusion cycles. From our analysis it follows that the slowest processes in the XCF cycle are the formation of positive ions ($^3\text{HedX}$) and ($^4\text{HedX}$). The corresponding formation times are estimated to be of the order of $10^{-8}$ s (see Appendix A1c). Therefore, for the $X$-particles to survive during $3.5 \times 10^9$ fusion cycles and produce $\sim 7 \times 10^4$ TeV of energy, their lifetime $\tau_X$ should exceed $\sim 10^8$ s. For shorter lifetimes the energy produced by a single $X$-particle before it gets stuck to a $^6\text{Li}$ nucleus is reduced accordingly.

### III. ACQUISITION AND REACTIVATION OF $X$-PARTICLES

The amount of energy produced by a single $X$-particle has to be compared with energy expenditures related to its production. $X$-particles can be produced in pairs in accelerator experiments, either in $l^+l^-$ annihilation at lepton colliders or through the Drell-Yan processes at hadronic machines. Although the energy $E \sim 7 \times 10^4$ TeV produced by one $X$-particle before it gets knocked out of the catalytic process is quite large on microscopic scale, it is only about 10 mJ. This means that $\gtrsim 10^8$ $X$-particles are needed to generate 1 MJ of energy. While colliders are better suited for discovery of new particles, for production of large numbers of $X$-particles fixed-target accelerator experiments are more appropriate. For such experiments the beam energy must exceed the mass of the $X$-particle significantly. Currently, plans for building such machines are being discussed [50].

The problem is, however, that the $X$-particle production cross section is very small. This comes about because of their expected large mass ($m_X \gtrsim 1\text{ TeV}/c^2$) and the fact that for their efficient moderation needed to make the formation of $(dX)$ atoms possible, $X$-particles should be produced with relatively low velocities. The cross section $\sigma_p$ of production of $X$-particles with mass $m_X \gtrsim 1\text{ TeV}/c^2$ and $\beta = v/c \approx 0.3$ is only $\sim 1$ fb (note that for scalar $X$-particles $\sigma_p \propto \beta^3$). As a result, the energy spent on production of an $X^{++}X^{--}$ pair will be by far larger than the energy that can be generated by one $X^{--}$ before it gets bound to a $^6\text{Li}$ nucleus. This means that reactivating and reusing the bound $X$-particles multiple times would be mandatory in this case. This, in turn, implies that only very long lived $X$-particles with $\tau_X \gtrsim 3 \times 10^4$ yr will be suitable for energy production.

Reactivation of $X$-particles bound to $^6\text{Li}$ requires dissociation of ($^6\text{LiX}$) ions. This could be achieved by irradiating them with particle beams, similarly to what was suggested for reactivation of lower-charge catalyst particles in ref. [46]. However, it would be much more efficient to use instead ($^6\text{LiX}$) ions as projectiles and irradiate a target with their beam. The Coulomb binding energy of $X$ to $^6\text{Li}$ is about 2 MeV; to strip them off by scattering on target nuclei with the average atomic number $A \approx 40$ one would have to accelerate ($^6\text{LiX}$) ions to velocities $\beta \approx 0.01$ which, for $m_X \approx 1\text{ TeV}/c^2$, corresponds to beam energy $\sim 0.05$ GeV. At these energies the cross section of the stripping reaction is $\gtrsim 0.1$ b, and $X$-particles can be liberated with high efficiency in relatively small targets. The energy spent on the reactivation of one $X$-particle will then only be about $10^{-9}$ of the energy it can produce before sticking to a $^6\text{Li}$ nucleus.

If $X$-particles are stable or practically stable, i.e. their lifetime $\tau_X$ is comparable to the age of the Universe, there may exist a terrestrial population of relic $X$-particles bound to nuclei or (in the case of $X^{++}$) to electrons and thus forming exotic nuclei or atoms. The possibility of the existence of exotic bound states containing charged massive particles was suggested in ref. [51] (see also [52]) and has been studied by many authors. The concentration of such exotic atoms on the Earth may be very low if reheating after inflation occurs at sufficiently low temperatures. Note that reheating temperatures as low as a few MeV are consistent with observations [53]. A number of searches for such superheavy exotic isotopes has been carried out using a variety of experimental techniques, and upper limits on their concentrations were established, see [54] for a review.

Exotic helium atoms ($X^{++}ee$) were searched for in the Earth’s atmosphere using laser spectroscopy technique, and the limit of their concentration $10^{-12} - 10^{-17}$ per atom over the mass range $20 - 10^4\text{ GeV}/c^2$ was established [55]. In the case of doubly negatively charged $X$, their Coulomb binding to nuclei of charge $Z$ would produce superheavy exotic isotopes with nuclear properties of the original nuclei but chemical properties of atoms with nuclear charge $Z - 2$. Such isotopes could have accumulated in continental crust and marine sediments. Singly positively charged ions ($^6\text{LiX}$) and ($^7\text{LiX}$) chemically behave as superheavy protons; they can capture electrons and form anomalously heavy hydrogen atoms. Experimental searches for anomalous hydrogen in normal water have put upper limits on its concentration at the level of $\sim 10^{-28} - 10^{-29}$ for the mass range 12 to 1200 $\text{GeV}/c^2$ [56] and $\sim 6 \times 10^{-15}$ for the masses between 10 and $10^3\text{ TeV}/c^2$ [57].

If superheavy isotopes containing relic $X$-particles of cosmological origin exist, they can be extracted from minerals e.g. by making use of mass spectrometry techniques, and their $X$-particles can then be stripped off.

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2 We thank M. Pospelov for this suggestion.
To estimate the required energy, we conservatively assume that it is twice the energy needed to vaporize the matter sample. As an example, it takes about 10 kJ to vaporize 1 g of granite [58]; denoting the concentration of X-particles in granite (number of X per molecule) by $c_X$, we find that the energy necessary to extract one X-particle is $\sim 2.3 \times 10^{-18} J/c_X$. Requiring that it does not exceed the energy one X-particle can produce before getting stuck to a $^3$Li nucleus leads to the constraint $c_X \gtrsim 2.3 \times 10^{-16}$. If it is satisfied, extracting X-particles from granite would allow XCF to produce more energy than it consumes, even without reactivation and recycling of the X-particles. Another advantage of the extraction of relic X-particles from minerals compared with their production at accelerators is that it could work even for X-particles with mass $m_X \gg 1$ TeV/c$^2$.

In assessing the viability of XCF as a mechanism of energy generation, in addition to pure energy considerations one should obviously address many technical issues related to its practical implementation, such as collection and moderation of the produced X-particles and prevention of their binding to the surrounding nuclei (or their liberation if such binding occurs), etc. However, the corresponding technical difficulties seem to be surmountable [59].

IV. DISCUSSION

There are several obvious ways in which our analysis of XCF can be generalized. Although we only considered nuclear fusion catalyzed by scalar X-particles, doubly charged particles of non-zero spin can do the job as well. While we studied XCF in deuterium, fusion processes with participation of other hydrogen isotopes can also be catalyzed by X-particles.

We considered XCF taking place in X-atomic states. The catalyzed fusion can also proceed through in-flight reactions occurring e.g. in $d + (dX)$ collisions. However, because even at the highest attainable densities the average distance $\bar{r}$ between deuterons is much larger than it is in $(ddX)$ atoms, the rates of in-flight reactions are suppressed by a factor of the order of $(\bar{r}/a_d)^3 \gtrsim 10^5$ compared with those of reactions occurring in X-atoms.

Our results depend sensitively on the properties of positive ions ($^{(4)}$HdX) and ($^{(3)}$HdX), for which we obtained only crude estimates. More accurate calculations of these properties and of the formation times of these positive ions would be highly desirable.

The existence of long-lived doubly charged particles may have important cosmological consequences. In particular, they may form exotic atoms, which have been discussed in connection with the dark matter problem [60–62]. They may also affect primordial nucleosynthesis in an important way. In ref. [63] it was suggested that singly negatively charged heavy metastable particles may catalyze nuclear fusion reactions at the nucleosynthesis era, possibly solving the cosmological lithium problem. The issue has been subsequently studied by many authors, see refs. [64, 65] for reviews. Doubly charged scalars X may also catalyze nuclear fusion reactions in the early Universe and thus may have significant impact on primordial nucleosynthesis. On the other hand, cosmology may provide important constraints on the XCF mechanism discussed here. Therefore, a comprehensive study of cosmological implications of the existence of $X^{\pm 2}$ particles would be of great interest.

To conclude, we have demonstrated that long-lived or stable doubly negatively charged scalar particles X, if exist, can catalyze nuclear fusion and provide a viable source of energy. Our study gives a strong additional motivation for continuing and extending the experimental searches for such particles.

Note added. Recently, the ATLAS Collaboration has reported a $3.6\sigma$ ($3.3\sigma$) local (global) excess of events with large specific ionization energy loss $|dE/dx|$ in their search for long-lived charged particles at LHC [88]. In the complete LHC Run 2 dataset, seven events were found for which the values of $|dE/dx|$ were in tension with the time-of-flight velocity measurements, assuming that the corresponding particles were of unit charge. It has been shown in [89] that this excess could be explained as being due to relatively long-lived doubly charged particles. It would be very interesting to see if the reported excess will survive with increasing statistics of the forthcoming LHC Run 3.

V. ACKNOWLEDGMENTS

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Appendix A: Atomic processes and X-atom formation times

1. Formation times of X-atomic systems

a. Formation of $(dX)$ ions and of $(ddX)$ and ($^{(3)}$HdX) atoms

Moderation of muons in medium and formation of $\mu$-atoms were considered in the classic papers [66, 67] which stood the test of time (see e.g. [68]). The moderation time is practically independent of the mass of the ionizing particle and is inversely proportional to square of its charge; this allows one to deduce the moderation times for X-particles by a simple scaling of the muonic case. From the results of ref. [67] we find that the moderation time of X-particles from $\beta \equiv v/c \simeq 0.1$ to atomic velocities $v \simeq 2c^2/\hbar \simeq 1.5 \times 10^{-2}c$ is

$$\tau \simeq 6 \times 10^{-11} \text{s} \quad (A1)$$
at liquid hydrogen density $N_0 = 4.25 \times 10^{22}$ nuclei/cm$^3$ and $T \approx 20$K. It is about $4.8 \times 10^{-8}$ s in deuterium gas at 0°C and pressure of one bar.

Once an X-particle has slowed down to atomic velocities, it gets captured on a highly excited state of the $(dX)$-ion, which then de-excites through a combination of $\gamma$-ray cascade emission (mostly E1 transitions) and inelastic scattering on the neighboring deuterium atoms with their Auger ionization. This is similar to de-excitation of highly excited $(\mu d)$ atoms in the case of $\mu$CF. In the latter case, the two de-excitations processes are generically of comparable rates; at liquid hydrogen density the Auger process slightly dominates. The de-excitation to the $(\mu d)$ ground state occurs within $t \sim 10^{-12}$ s [69].

In the case of $(dX)$ de-excitation, the radiative processes get enhanced. Indeed, the rates of E1 emission are proportional to cube of the energy $E_\gamma$ of the emitted $\gamma$ and square of the transition matrix element of the electric dipole operator $d_{fi}$. It is easy to see that $E_\gamma$ scales linearly with the mass of the atomic orbiting particle $m$, whereas $d_{fi}$ scales linearly with the Bohr radius of the system, i.e. is inversely proportional to $m$. Therefore, the rates of E1 transitions scale linearly with $m$. As a result, we find that the rate of radiative de-excitation of the $(X d)$ system is larger than that of $(d\mu)$ atom by a factor $m_{d\mu}/m_\mu \sim 20$. One can therefore expect the de-excitation of $(dX)$ ions to be at least as fast as that for the muonic deuterium, i.e. to occur within $\sim 10^{-12}$ s.

The produced negative ion $(dX)$ can pick up another deuterium from the medium to form a highly excited state of the $(d(dX))$ atom, which can de-excite through the same processes as $(dX)$. In addition, being electrically neutral, $(d(dX))$ can penetrate deep inside the neighboring deuterium atoms and experience the electric field of their nuclei. This leads to Stark mixing effects which further accelerate the de-excitation processes [69, 70].

The situation is quite similar for the formation time of $(3^3HeX)$ atoms. An $(3^3HX)$ negative ion produced in reaction (2b) picks up a deuterium from the medium to form a highly excited state of $(3^3HeX)$ atom. The latter then de-excites as described above for the $(d(dX))$ atom, within approximately the same time interval.

b. Charge exchange reaction $d+(pX) \rightarrow (dX)+p$

A simple and accurate estimate of the cross section of the muon exchange reaction $d+(\mu d) \rightarrow (d\mu)+p$, based on dimensional analysis and the fact that low-energy cross sections of inelastic processes are inversely proportional to the relative velocities of the colliding particles, was given in [36]. The cross section of the reaction $d+(pX) \rightarrow (dX)+p$ can be estimated similarly, which yields

$$\sigma_e \simeq 4\pi a_p^2 f v_e/v.$$  \hspace{1cm} (A2)

Here $a_p = \hbar/(2\alpha m_p c) = 1.44 \times 10^{-12}$ cm is the Bohr radius of $(pX)$ atom, $v$ and $v_e$ are the relative velocities of the involved particles in the initial and final states, respectively, and $f$ is a constant of order unity. Taking into account that the relative velocities of the initial-state particles are very small and the $Q$-value of the reaction $d+(pX) \rightarrow (dX)+p$ is $\approx 90$ keV, we find $v_e \approx 1.4 \times 10^{-2} c$, which gives

$$\sigma_e v \approx 10^{-14} \text{ cm}^2/s.$$  \hspace{1cm} (A3)

For the rate $\lambda_e$ and the characteristic time $t_c$ of this reaction we then find, at liquid hydrogen density $N_0 = 4.25 \times 10^{22}$ nuclei/cm$^3$,

$$\lambda_e = \sigma_e v N_0 \approx 4 \times 10^8 \text{ s}^{-1}, \quad t_c = \lambda_e^{-1} \approx 2.5 \times 10^{-9} \text{ s}.$$  \hspace{1cm} (A4)

c. Positive ions $(3^3HeX)$ and $(4^4HeX)$ and time scales of their formation

The complexes $(3^3HeX)$ and $(4^4HeX)$ are positive ions. Such systems can be considered as composed of a tightly bound “inner core”, represented by a neutral $(HeX)$ atom, and a deuteron, weakly bound to the core by atomic polarization effects. The neutral atoms in the inner cores are slightly perturbed by the presence of an external deuteron and are characterized by the binding energies and Bohr radii approximately equal to those of the corresponding $(3^3He)$ or $(4^4He)$ atoms in the absence of the additional deuteron. The extraneous deuteron is bound on a $1s'$ orbit characterized by a larger radius and much smaller binding energy.

It is not immediately obvious if such exotic atomic systems are actually stable; in particular, their stability cannot be deduced from the stability of negative ion of hydrogen $H^−$ familiar from the usual atomic physics. Stability of three-body Coulomb systems with arbitrary masses and charges of the particles was studied in a number of papers, see e.g. [47–49]. From their general results it follows that the states $(3^3HeX)$ and $(4^4HeX)$ should actually be stable. This can be seen from fig. 8 of ref. [47], fig. 3 of [48] or fig. 13 of [49], where the stability regions are shown in the case of arbitrary fixed masses of the particles and the charge $q_1 = 1$ as a function of $q_2$ and $q_3$. Here $q_1$ is the absolute value of the charge of the particle which is opposite to the other two ($q_1 = Z_{Xe}$ in the case we consider), whereas $q_2$ and $q_3$ are the charges of the same-sign particles, with the convention $q_2 \geq q_3$. As the stability depends on the ratios of the charges and not on their absolute values, $q_1$ was set equal to unity for convenience. With such a normalization, we have $q_2 = 1$, $q_3 = 1/2$. It can be seen from the above mentioned figures that the point $(q_2^{-1}, q_3^{-1}) = (1, 2)$ is inside the stability region, that is, positive ions $(3^3HeX)$ and $(4^4HeX)$ must be stable. This point is, however, rather close to the border of the stability region, which reflects the relative smallness of the binding energy of the deuteron (“deuteron affinity”).

We have attempted a variational calculation of the binding energies of such positive ions using simple two-
and three-parameter Hylleraas-type trial wave functions which were able to predict the stability of $H^-$ ion, but found no binding of deuteron. This is apparently related to the fact that $(\text{Hed}X)$ ions, which have nuclei of differing mass and charge on their atomic orbits, are more complex than $H^-$ ion, whose two electrons are identical particles. A qualitative analysis of the properties of such systems would therefore necessitate calculations with more sophisticated trial wave functions. This would require a dedicated study, which is beyond the scope of the present paper.

In the absence of an actual calculation, we have to resort to semi-quantitative methods. In doing that, we will be using the properties of negative ion $H^-$ as a starting point, but will also take into account the peculiarities of $(\text{Hed}X)$ and $(\text{Hed}X)$ ions. In the case of $H^-$, the radius of the outer electron’s orbit is about a factor 3.7 larger than that of the inner electron [71], and the binding energy of the outer electron (electron affinity) is about 18 times smaller than that of the inner one. Taking into account tighter binding of the inner core in the case of the positive ions we consider, we assume the radii $a$ of their external orbits and the deuteron binding energies $E_{bd}$ to be, respectively, a factor of $\sim 30$ larger and three orders of magnitude smaller than those of the corresponding (He$^X$) atoms. Factor $\sim 30$ increase for $a$ compared with the inner core radius is obtained as follows: we multiply the factor 3.7 hinted by $H^-$ ions by $2 \times 2 = 4$ due to the $X$ particle and the He nucleus each having charge 2 and by the ratio of the mass of $^3\text{He}$ (or $^4\text{He}$) to the deuteron mass. This gives factor $\sim 22 - 30$. For the binding energy of the extra deuteron we take into account that it scales as $a^{-2}$. We therefore choose

$$(\text{Hed}X): \quad a \simeq 7 \times 10^{-12} \text{cm}, \quad E_{bd} \simeq 1.2 \text{keV}, \quad (A5)$$

$$(\text{Hed}X): \quad a \simeq 5 \times 10^{-12} \text{cm}, \quad E_{bd} \simeq 1.6 \text{keV}. \quad (A6)$$

The formation of $(\text{Hed}X)$ and $(\text{Hed}X)$ ions can proceed as follows. An $(\text{He}^X)$ atom produced in reaction (1b) collides with the neighboring $D_2$ molecules, dissociating them and picking up one of their deuterons through the exothermic reaction

$$(\text{He}^X) + D_2 \rightarrow (\text{Hed}X) + d + 2e^- . \quad (A7)$$

This is the dissociative attachment (DA) mechanism, analogous to the one by which $H^-$ ions are produced in $e^- + \text{H}_2 \rightarrow \text{H}^- + \text{H}$ reactions. An important difference is, however, that what is attached is now a nucleus (deuteron) rather than an electron. The formation of $(^4\text{He}X)$ ions from $(^3\text{He}X)$ atoms produced in reaction (4b) and (5b) proceeds similarly. [Note that a tiny fraction of $(^4\text{He}X)$ ions is produced directly in stage I reaction (3b)].

As the $Q$-values of the formation reactions of $(^3\text{He}X)$ and $(^4\text{He}X)$ ions are about two orders of magnitude larger than the dissociation energy of $D_2$ molecules and the ionization potential of $D$ atoms, these processes are actually similar to the usual charge exchange reactions on free particles, except that most of the released energy is now carried away by the final-state electrons. The rates and characteristic times of these processes can therefore be estimated using the expressions similar to those in eqs. (A3) and (A4). This gives, at the liquid hydrogen density,

$$\lambda_{DA} \sim 5 \times 10^7 \text{s}^{-1}, \quad t_{DA} = \lambda_{DA}^{-1} \sim 2 \times 10^{-8} \text{s} . \quad (A8)$$

2. Atomic binding energies of light nuclei in $X$-atoms and $Q$-values of fusion reactions

The $Q$-value of an XCF reaction can be found by subtracting from the $Q$-value of the corresponding $X$-less reaction the atomic binding energy of the nuclei in the initial state and adding to it the binding energy of one of the produced nuclei to $X$ in the final state, when such bound states are formed. We therefore first find the relevant binding energies.

a. Antihelium-like $(ddX)$ atom

The total binding energy of helium atom is 79.005 eV. From this value, the binding energy of $(ddX)$ atom is obtained to a very good accuracy by the simple rescaling with the factor $m_d/m_e$, which gives $E_b(ddX) = 0.290 \text{ MeV}$.

b. Other $X$-atoms with Coulomb-bound light nuclei

For atomic $(NN'X)$ states other than $(ddX)$ we first consider hydrogen-like atoms $(NX)$ and then estimate the atomic binding energy of the additional nucleus $N'$ ($m > m'$ is assumed). In the limit of pointlike nuclei the ground-state wave function $\psi_{1s}(r)$, the Bohr radius $a$ and the binding energy $E^0_b$ of an $(NX)$ state are

$$\psi_{1s}(r) = \frac{1}{\sqrt{\pi a^3}} e^{-r/a}, \quad a = \frac{\hbar^2}{Z_XZe^2m} = \frac{\hbar}{Z_XZe^2mc}, \quad E^0_b = |E^0_{1s}| = \frac{1}{2} (Z_XZa)^2mc^2 . \quad (A9)$$

Here $Ze$ and $m$ are the charge and the mass of the nucleus $N$, and $-Z_Xe$ is the charge of the $X$-particle ($Z_X = 2$ in
the case under discussion).

For most of the nuclei we consider, Bohr radii of the \((NX)\) atomic states are either comparable to or smaller than the nuclear radii, and the approximation of pointlike nuclei is rather poor. We therefore allow for finite nuclear sizes by making use of a variational approach. We consider nuclei as uniformly charged balls of radius \(R\) and employ the simple one-parameter test wave function of Flügge and Zickendrant, which has the correct asymptotics for both large and small \(r\) [72, 73]:

\[
\psi(r) = N(\lambda) \left(1 + \frac{\lambda r}{2R}\right) e^{-\lambda r}, \quad N(\lambda) = \sqrt{\frac{1}{\pi} \left(\frac{\lambda}{2R}\right)^3}.
\] (A10)

Here \(\lambda\) is the variational parameter. With this wave function, the expectation value of the energy of the system is

\[
E(\lambda) = \frac{3}{56} \frac{\hbar^2}{mR^2} \left\{ \lambda^2 + \frac{R}{a} \left[ \frac{216}{\lambda^2} - 28 - e^{-\lambda} \left( \frac{216}{\lambda^2} + \frac{216}{\lambda} + 80 + 14\lambda + \lambda^2 \right) \right] \right\}.
\] (A11)

We minimize it numerically, which yields the ground-state energy \(E_{1s}\), the binding energy of the system being \(E_b(R) = |E_{1s}|\). The corresponding value of \(\lambda\) determines, through eq. (A10), the ground-state wave function of the system. It will be used in the calculations of the sticking probabilities in sec. 3.3a below.

We calculate the binding energies of \((NX)\) states for two different choices of the values of the nuclear radii \(R\). First, we employ the frequently used expression \(R = R_N \equiv 1.2A^{1/3}\) fm, where \(A\) is the atomic number of the nucleus \(N\). Second, we make use of the experimentally measured rms charge radii \(r_{NC} \equiv \langle r_c^2 \rangle^{1/2}_N\) [74] and set the nuclear radii equal to \(R_{NC} \equiv (5/3)^{1/2} r_{NC}\), which is the relation between \(R_{NC}\) and \(r_{NC}\) in the uniformly charged ball model of the nucleus. The results are presented in Table I along with the Bohr radii \(a\) and the binding energies for pointlike nuclei \(E_b^0\). Note that for our further calculations we use the results based on the experimentally measured nuclear charge radii, which are presumably more accurate.

As a test, we also performed similar calculations for atomic systems \((NC)\) with \(C\) a singly negatively charged heavy particle, for which the binding energies were previously found in [63]. Our results are in good agreement with those of ref. [63], the difference typically being within 1%.

The binding energies of the \(X\)-atoms in the initial states of the XCF reactions other than \((ddX)\) are found as follows. For \((^3\text{He}dX)\) atoms, we add to the binding energy of negative \((^3\text{HX})\) ion the deuteron binding energy found through the variational procedure described above, assuming deuteron to be a pointlike particle in the Coulomb field of a nucleus of charge \(Z_{\text{He}} - Z_X = -1\) and radius equal to that of \(^3\text{H}\) nucleus. For positive ions \((^3\text{He}dX)\) and \((^4\text{He}dX)\), we add to the binding energies of \((^3\text{He}X)\) and \((^4\text{He}X)\) atoms the deuteron binding energy \(E_{bd}\) given in eqs. (A5) and (A6), respectively. The obtained binding energies are then used for calculating the \(Q\)-values of the XCF reactions under consideration. The results are shown in the third column of Table III.

3. Sticking probabilities and related issues

a. Sticking probabilities in the sudden approximation

To evaluate the probability \(\omega_s\) that the \(X\)-particle in the final state of a fusion process will stick to one of the produced nuclear fragments we make use of the fact that nuclear reactions of XCF occur on time scales that are much shorter than the characteristic \(X\)-atomic time. Indeed, the characteristic time of \(X\)-atomic processes is \(t_{at} \sim a_d/v_{at} \sim \hbar^3/(4m_e^2e^4) \approx 1.6 \times 10^{-21}\) s, whereas the fusion reactions of XCF occur on the nuclear time scales \(\lesssim 10^{-23}\) s. This disparity between the atomic and nuclear time scales in XCF allows one to use the sudden approximation [75] for evaluating the \(X\)-sticking probabilities \(\omega_s\).

Consider the reaction \((N_1N_2X) \rightarrow N_3 \pm N_4 + X\). Just before the fusion occurs, the nuclei in the \((N_1N_2X)\) atom approach each other to a distance of the order of the range of nuclear force. The atomic wave function therefore adiabatically goes over into that of the hydrogen-like atom with a nucleus of mass \(m_1 \pm m_2\) and charge \(Z_1 = Z_i + Z_2\) orbiting the \(X\) particle. We denote this wave function \(\psi_i\). As the fusion occurs suddenly (compared with the atomic time scale), the state of the atomic system immediately after the fusion will be described by the same wave function. Transition amplitudes can then be found by projecting it onto the proper final states. Let the velocity of the produced nucleus \(N_3\) of mass \(m_3\) be \(\vec{v}_3\). As the final-state \(X\)-particle is practically at rest, this is also the relative velocity of \(N_3\) and \(X\). The probability that \(N_3\) will get captured by \(X\) and form a bound state with it is then

\[
\omega_s = \sum_\alpha \left| \int \psi^{*}_{fa} \psi_i e^{-i \vec{q} \vec{r} \cdot dV} \right|^2,
\] (A12)

where \(\psi_{fa}\) is the wave function of the final \((N_3X)\) state and \(\vec{q} = m_3 \vec{v}_3/h, m_3 = m_3m_X/(m_3 + m_X) \approx m_3\) being the reduced mass of the \(N_3X\) system. The sum in (A12) is over all the bound states of hydrogen-like atom \((N_3X)\). The case of radiative fusion reactions \((N_1N_2X) \rightarrow \)
| Bound state | Bohr radius $a$ (fm) | $r_{Nc}$ (fm) [74] | $R_N = 1.2A^{1/3}$ (fm) | $R_{Nc}$ (fm) | $E_b(R_N)$ (MeV) | $E_b(R_{Nc})$ (MeV) | $E_b^0$ (MeV) |
|------------|----------------------|------------------|-----------------|-------------|-----------------|-----------------|-----------|
| ($pX$)     | 14.4                 | 0.8783           | 1.20            | 1.134       | 0.096           | 0.096           | 0.100     |
| ($dX$)     | 7.20                 | 2.142            | 1.51            | 2.765       | 0.189           | 0.183           | 0.200     |
| ($^3HX$)   | 4.81                 | 1.759            | 1.73            | 2.271       | 0.276           | 0.268           | 0.299     |
| ($^3HeX$)  | 2.41                 | 1.966            | 1.73            | 2.538       | 1.00            | 0.905           | 1.196     |
| ($^4HeX$)  | 1.81                 | 1.676            | 1.905           | 2.163       | 1.202           | 1.153           | 1.588     |
| ($^6LiX$)  | 0.805                | 2.589            | 2.18            | 3.342       | 2.680           | 2.069           | 5.369     |

Table I. Properties of ($NX$) bound states. Third and fifth columns show experimental values of rms charge radii $r_{Nc} \equiv \langle r_{c,N}^2 \rangle^{1/2}$ from ref. [74] and the corresponding nuclear radii found as $R_{Nc} = (5/3)^{1/2}r_{Nc}$. $E_b(R_N)$ and $E_b(R_{Nc})$ are binding energies calculated for the corresponding values of nuclear radii; $E_b^0 = (Z_N Z_c)^2 me^2/2$ is binding energy in the limit of pointlike nuclei.

$N_3 + \gamma + X$ is considered quite similarly. However, due to different kinematics, the values of $q = |\vec{q}|$ are related differently to the $Q$-values for these reactions. For the non-radiative reactions we have $q = \sqrt{2\mu Q}/h$ with $\mu = m_3 m_a/(m_3 + m_a)$, whereas for the radiative ones $q \approx Q/(\hbar c)$.

The main contribution to $\omega_s$ comes from the transition to the ground state of ($N_3 X$) atom, with total contribution of all the excited states being less than 20% [34]. For our estimates we shall therefore restrict ourselves to transitions to the ground states. The functions $\psi_i$ and $\psi_f$ are then the wave functions of the $1s$ states of the hydrogen-like atoms with masses and charges of the atomic particles $m_i = m_1 + m_2$, $Z_i = Z_1 + Z_2$ and $m_f = m_3$, $Z_f = Z_3$, respectively.

To take into account the finite size of the nuclei, we use the wave functions (A10) with the substitutions $\lambda \rightarrow \lambda_{i,f}$, where the variational parameters $\lambda_{i,f}$ are found from the minimization of $E(\lambda)$ defined in (A11) with the replacements $a \rightarrow a_{i,f}$ and $R \rightarrow R_{i,f}$. The Bohr radii $a_{i,f}$ are given by the standard formula (see eq. (A17) below); the nuclear radii $R_{i,f}$ can be found from the rms nuclear charge radii as discussed in sec. A2b. To find $R_{i,f}$, we can directly use the experimentally measured rms charge radius of the nucleus $N_3$. For the initial state, we approximate the rms charge radius $r_i$ of the compound nucleus $N_1 N_2$ as

$$r_i \approx (r_{N1c}^3 + r_{N2c}^3)^{1/3}, \quad \text{(A13)}$$

where $r_{N1c} \equiv \langle r_{c,N1}^2 \rangle^{1/2}$ and $r_{N2c} \equiv \langle r_{c,N2}^2 \rangle^{1/2}$ are the experimentally measured rms charge radii of the $N_1$ and $N_2$ nuclei, respectively. Note that eq. (A13) corresponds to the liquid drop model of nucleus. Eq. (A12) then gives for the sticking probability

$$\omega_s = \left[ \frac{32\pi N_i \kappa}{(\kappa^2 + q_i^2)^3} \left( \kappa^2 + \frac{3\kappa \kappa_i (\kappa_i^2 - q_i^2)}{\kappa^2 + q_i^2} \right) \right]^2, \quad \text{(A14)}$$

where $N_{i,f} = N(\lambda_{i,f})$, $\kappa_{i,f} = \lambda_{i,f}/2R_{i,f}$, $\kappa = \kappa_i + \kappa_f$. \text{(A15)}

For comparison, we also calculate the sticking probabilities $\omega_{s0}$ neglecting the nuclear size, i.e. employing the usual ground-state wave functions of the hydrogen-like atoms with pointlike nuclei (A9). From eq. (A12) we find

$$\omega_{s0} = \frac{(2a_r)^6}{(a_{i,f} a_r)^3 (1 + q^2 a_r^2)^4}, \quad \text{(A16)}$$

where

$$a_{i,f} = \frac{\hbar}{Z_X Z_f c m_i c}, \quad a_r = \frac{a_{i,f}}{a_{i} + a_{f}}. \quad \text{(A17)}$$

The obtained values of the sticking probabilities $\omega_s$ and $\omega_{s0}$ are shown in the fourth and fifth columns of Table III.

b. Lifting the prohibition of radiative E1 transitions for Coulomb-bound nuclei

The radiative nuclear fusion reaction $d + d \rightarrow ^4He + \gamma$ has a tiny branching ratio. This is because it proceeds mostly through E2 electromagnetic transitions, as E1 transitions are strictly forbidden for fusions of identical particles. Indeed, after the separation of the irrelevant center-of-mass motion, one finds that for particles of charges $q_{1,2}$ and masses $m_{1,2}$ the effective charge of the electric dipole operator is $q = (q_1 m_2 - q_2 m_1)/(m_1 + m_2)$, which vanishes when the two particles have the same charge-to-mass ratio.

The situation may be different for XCF, when the fusion particles are bound to an orbit of an X-atom. Fusion may then proceed through the transition to an intermediate excited atomic state, which then de-excites via atomic electric dipole radiation. In this mechanism the motion of the center of mass of the initial-state nuclei plays a central role, and the effective charge of the dipole operator does not vanish. Consider the XCF reaction

$$(ddX) \rightarrow (^4HeX)^* \rightarrow (^4HeX) + \gamma, \quad \text{(A18)}$$
where \((ddX)\) is in its ground \((1s)^2\) state and \((^4\text{He}X)^*\) is an excited state of \((^4\text{He}X)\) atom which can decay through an E1 transition. For definiteness, we take \((^4\text{He}X)^*\) to be a state with the principal quantum number \(n = 3\) (contributions of states with higher \(n\) are in general suppressed as \(1/n^3\)). Conservation of the total angular momentum and parity in strong interactions responsible for the fusion process imply that the intermediate state \(n = 3\) can be either \(3s\) or \(3d\). The excited \((^4\text{He}X)^*\) state can then decay through E1 \(\gamma\)-emission to the \(2p\) state of \((^4\text{He}X)\) atom, which will eventually de-excite to the ground state of \((^4\text{He}X)\). Thus, the radiative fusion reaction (3b) might proceed through an E1 transition from an excited state of \((^4\text{He}X)\) rather than through the usual E2 transitions.

It is easy to see, however, that this does not lead to any appreciable increase of the rate of reaction (3b). Indeed, the amplitude of the process in eq. (A18) contains the product of the amplitude of fusion with the formation of \((^4\text{He}X)^*\) and the amplitude of its subsequent E1 de-excitation. The rate of the process (A18) is therefore proportional to the probability that \(^4\text{He}\) produced as a result of the fusion reaction is bound to \(X\) in an excited state of the \((^4\text{He}X)\) atom rather than being ejected. The latter can be found in the sudden approximation by making use of the expression on the right-hand side of eq. (A12) with \(\psi_f^a\) being the wave functions of the 3s and 3d states. Using for our estimates the hydrogen-like wave functions for pointlike nuclei, we find for the corresponding probabilities

\[
P_{3s} = \frac{2^8 (q^2a^2)^2(q^2a^2 + \frac{16}{9})^2}{(q^2a^2 + \frac{16}{9})^3}, \quad P_{3d} = \frac{2^{17}}{3^9} (q^2a^2 + \frac{16}{9})^8.
\]

Here \(q \approx Q/(\hbar c)\) is the momentum transfer to the produced \(^4\text{He}\) nucleus divided by \(\hbar\), and \(a\) is the Bohr radius of \((^4\text{He}X)\) atom. The rate of the process (A18) is therefore proportional to the probability that \(^4\text{He}\) produced as a result of the fusion reaction is bound to \(X\) in an excited state of the \((^4\text{He}X)\) atom rather than being ejected. The latter can be found in the sudden approximation by making use of the expression on the right-hand side of eq. (A12) with \(\psi_f^a\) being the wave functions of the 3s and 3d states. Using for our estimates the hydrogen-like wave functions for pointlike nuclei, we find for the corresponding probabilities

Here \(R\) is the nuclear radius \(R_N\) for nuclear transitions and the size of the atomic system \(a\) for transitions between atomic states. Note that, in the case of \(X\)-atoms with light nuclei, the nuclear and atomic radii are of the same order of magnitude (see Table 1). For the process under consideration one could expect, neglecting the suppression of E1 transitions, \(\Gamma(E1)_{\text{unsup}}/\Gamma(E2) \sim (625/12)(E_e/a/\hbar c)^{-2} \sim 800\). Taking into account the discussed above suppression factor, we arrive at \(\Gamma(E1)/\Gamma(E2) \sim 1.4 \times 10^{-3}\).

Thus, although transitions through excited atomic states of \((^4\text{He}X)\) atom lift the forbiddance of electric dipole radiation in the fusion reaction (3b), the resulting E1 transition is heavily hindered, and its contribution to the rate of the process can be neglected.

A very similar argument applies to the radiative fusion reaction (6b). Although the fusing nuclei, \(^4\text{He}\) and \(d\), are not identical in this case, they have nearly the same charge-to-mass ratio. As a result, in the corresponding \(X\)-less reaction E1 transitions are heavily suppressed, and the reaction proceeds primarily through E2 radiation. The XCF reaction (6b) could go through an excited atomic states of \((^6\text{Li}X)\) atom. We find, however, that in this case the suppression of the atomic E1 transition is even stronger than it is for reaction (3b). Indeed, the \(Q\)-value of the reaction is rather small \((Q \approx 2.4\text{MeV})\), leading to \(qa \approx 10^{-2}\). Assuming again transitions through \(n = 3\) atomic states, we find for the probabilities of formation of the excited states \((^6\text{Li}X)^*\) the values \(P_{3s} \approx 3 \times 10^{-10}\), \(P_{3d} \approx 6 \times 10^{-10}\). The suppression factor due to the non-resonant nature of the radiative transitions from the 3s and 3d states to the \(2p\) state of \((^6\text{Li}X)\) is \((\Delta E/E_e)^2 \approx 0.02\). Altogether, this gives for the atomic E1 transitions in process (6b) the suppression factor \(\sim 2 \times 10^{-11}\), which makes them completely irrelevant.

**Appendix B: Astrophysical S-factors, reaction factors and fusion rates**

In this section we give the details of our calculations of the cross sections, rates and branching ratios of the XCF reactions under discussion.

1. **Cross sections and reaction factors**

The cross section of a fusion reaction of nuclei \(N_1\) and \(N_2\) of masses \(m_{1,2}\) and charges \(Z_{1,2}\) is usually written as [77]

\[
\sigma(E) = S(E) = \frac{S}{E} e^{-2\pi q_{12}}, \quad (B1)
\]
where $S(E)$ is the so-called astrophysical factor, $E$ is the c.m.s. energy and $\eta_{12}$ is the Sommerfeld parameter:

$$\eta_{12} = \frac{Z_1 Z_2 e^2}{\hbar v} = Z_1 Z_2 \alpha \sqrt{\frac{\mu c^2}{2E}}, \quad \mu = \frac{m_1 m_2}{m_1 + m_2}. \quad (B2)$$

Here $v$ is the relative velocity of the fusing particles. If there are no low-energy resonances in the fusion reaction, the astrophysical factor $S(E)$ is a slowly varying function of $E$ at low energies. For catalyzed fusion from the relative $s$-wave state of $N_1$ and $N_2$ the reaction factor $A(E)$ is defined as $A(E) = \sigma(E) v C_0^{-2}$ [32]. Here $C_0^2$ is the $s$-wave Coulomb barrier penetration probability factor:

$$C_0^2 = \frac{2\pi \eta_{12}}{e^{2\pi \eta_{12}} - 1}. \quad (B3)$$

This gives

$$A(E) = \frac{S(E)}{\pi Z_1 Z_2 \alpha \mu c} \left( 1 - e^{-2\pi \eta_{12}} \right). \quad (B4)$$

The transition from $S(E)$ to $A(E)$ takes into account the fact that the catalyst particle screens the Coulomb fields of the fusing nuclei and essentially eliminates the Coulomb barrier. The rates $\lambda$ of XCF reactions are related to the corresponding $A(E)$-factors as

$$\lambda = A(E) \rho_0, \quad \rho_0 = |\psi_i(0)|^2. \quad (B5)$$

Here $\rho_0$ is the squared modulus of the atomic wave function of the initial $(N_1 N_2 X)$ state taken at zero separation between the nuclei $N_1$ and $N_2$ (more precisely, at a distance of the order of the range of nuclear forces) and integrated over their distance $R$ to the $X$-particle. It plays the same role as the number density $n$ of the target particles in the usual expression for the reaction rates $\lambda = \sigma n v$ [32].

In muneic molecules or molecular ions, the energies of relative motion of nuclei are very low, so that $\eta_{12} \gg 1$; the term $e^{-2\pi \eta_{12}}$ in eq. (B4) is therefore always omitted in the literature on $\mu$CF. In addition, in evaluating the cross sections of the fusion reactions it is usually sufficient to consider the astrophysical $S$-factors in the limit $E \to 0$.

In contrast to this, in XCF processes the kinetic energy $E$ of the relative motion of the nuclei $N_1$ and $N_2$ in $(X N_1 N_2)$ atoms is not negligible. For a system of charges bound by the Coulomb force, the virial theorem relates the mean kinetic energy $T$ and mean potential energy $U$ as $2T = -U$. Therefore, the mean kinetic energy $T$ in the ground state coincides with the binding energy of the system: $E_b = |T + U| = T$. On the other hand, $T$ is the sum of the mean kinetic energies of the relative motion of $N_1$ and $N_2$ and of their center-of-mass motion.

This can be explained as follows. Consider for simplicity the wave function of $(N_1 N_2 X)$ atom to be a product of the hydrogen-like wave functions of $(N_1 X)$ and $(N_2 X)$ systems, i.e.

$$\Psi_i(\vec{r_1}, \vec{r_2}) = \frac{1}{\pi (a_1 a_2)^{3/2}} e^{-\frac{r_1^2}{a_1^2} - \frac{r_2^2}{a_2^2}}. \quad (B6)$$

Such a wave function obtains when one neglects the Coulomb interaction between $N_1$ and $N_2$, but in fact $\Psi_i$ of this form can partly include correlations between $N_1$ and $N_2$ provided that one replaces the charge of the $X$-particle $Z_X$ by an effective charge $Z_{X_{\text{eff}}}$ in the expressions for the Bohr radii $a_1$ and $a_2$ or, better still, treats $a_1$ and $a_2$ as variational parameters. The ground-state mean values of the total kinetic energy of the $(N_1 N_2 X)$ system, kinetic energy of the center of mass of $N_1$ and $N_2$ and their relative kinetic energy are then

$$\bar{T} = \frac{\hbar^2}{2m_1 a_1^2} + \frac{\hbar^2}{2m_2 a_2^2}, \quad \bar{T}_{\text{c.m.}} = \frac{\hbar^2}{2m} \left( \frac{1}{a_1^2} + \frac{1}{a_2^2} \right), \quad \bar{T}_{\text{rel}} \equiv E = \frac{\hbar^2}{2m_1 a_1^2} \frac{m_2}{m} + \frac{\hbar^2}{2m_2 a_2^2} \frac{m_1}{m}, \quad (B7)$$

where $m \equiv m_1 + m_2$. Note that $\bar{T}_{\text{c.m.}} + \bar{T}_{\text{rel}} = \bar{T}$. For $m_1 = m_2$ we find $E = \bar{T}/2 = E_b/2$. Assume now $m_1 > m_2$. Since we expect $a_i \propto m_i^{-1/2}$ $(i = 1, 2)$, the kinetic energies in eq. (B7) can be estimated as

$$E = \bar{E} - \bar{T}_{\text{c.m.}} \sim \frac{\hbar^2}{2m_1 a_1^2} \frac{m_2}{m} \sim \frac{\bar{T}_{\text{rel}}}{m}, \quad (B8)$$

Wave function (B6) can also be used for evaluating the
parameter $\rho_0$ defined in eq. (B5). Direct calculation gives

$$\rho_0 = \frac{1}{\pi(a_1 + a_2)^2}. \quad (B9)$$

We shall also employ the wave function $\Psi_i(r_1, r_2)$ of eq. (B6) in sec. B 3 for evaluating of the parameter $\rho_1$ that enters into the expression for the rates of reactions (3c) and (6c).

2. Astrophysical $S$-factors and XCF reaction rates

The Coulomb binding of the fusing nuclei $N_1$ and $N_2$ to an X-particle should have no effect on the strong interactions responsible for the fusion and can only modify the reaction rates due to the facts that (i) the Coulomb repulsion barrier is actually eliminated due to the very close distance between $N_1$ and $N_2$ in the X-atom, and (ii) because of the very small size of X-atoms, the number densities of $N_1$ and $N_2$ within an $(N_1N_2 X)$-atom are many orders of magnitude larger than their number densities achievable for in-flight fusion. This means that, for those reactions that can occur in the absence of X-particles, we can use the experimentally measured values of the corresponding astrophysical $S$-factors in order to calculate the rates of the XCF reactions. We take the relevant data from refs. [78–83]. For internal conversion (IC) reactions (3c) and (6c), which do not have relevant data from refs. [78–83]. For internal conversion (IC) reactions (3c) and (6c), the matrix element of an $E_0$ transition responsible for the fusion and can only modify the reaction rates due to the facts that (i) the Coulomb binding of the fusing nuclei $N_1$ and $N_2$ to an X-particle should have no effect on the strong interactions responsible for the fusion and can only modify the reaction rates due to the facts that (ii) because of the very small size of X-atoms, the number densities of $N_1$ and $N_2$ within an $(N_1N_2 X)$-atom are many orders of magnitude larger than their number densities achievable for in-flight fusion. This means that, for those reactions that can occur in the absence of X-particles, we can use the experimentally measured values of the corresponding astrophysical S-factors in order to calculate the rates of the XCF reactions. We take the relevant data from refs. [78–83]. For internal conversion (IC) reactions (3c) and (6c), which do not have X-less analogues, we calculate the rates directly in the next subsection. The input data necessary for the calculations of the reaction factors $A(E)$ for all the discussed reactions and the obtained results are shown in Table II.

3. Rates of internal conversion (IC) processes (3c) and (6c)

IC is de-excitation of an excited nucleus in an atom through the ejection of an atomic electron (or, in the case of muonic atoms, of a muon) [84]. For IC in a process of nuclear fusion, the initial excited nuclear state is a compound nucleus formed by the merger of the two fusing nuclei. In the case of $\mu$CF, the ejected particle is the muon; for XCF, it is more appropriate to speak about the ejection of the final-state nucleus itself rather than of the X-particle, as the latter is expected to be much heavier than light nuclei. However, when considered in terms of the relative motion between the “nucleus” and the orbiting particle, the treatment of IC in XCF closely parallels that in the case of the usual atoms or molecules.

At low energies relevant to fusion of light nuclei, IC predominantly proceeds through electric monopole (E0) transitions whenever this is allowed by angular momentum and parity selection rules. This is the case for reactions (3c) and (6c). The matrix element of an E0 transition can be written as [85]

$$M_{fi} = \frac{2\pi}{3} Z_X e^2 \bar{Q}_0 \psi_f^*(0) \psi_i(0). \quad (B10)$$

Here $\psi_i(0)$ is the atomic wave function of the initial-state compound nucleus bound to the X-particle and $\psi_f(0)$ is the final-state continuum atomic wave function of the ejected nucleus, both taken at zero separation between the nucleus and the X particle. The quantity $\bar{Q}_0$ (not to be confused with $Q_0$ of table III) is the transition matrix element of the nuclear charge radius operator between the initial and final nuclear states:

$$\bar{Q}_0 = \langle f | \sum_{i=1}^{Z} r_{pi}^2 | i \rangle. \quad (B11)$$

Here the sum is taken over nuclear protons. The rate of the process is readily found from the matrix element (B10):

$$\lambda_{IC} = g_s \sqrt{\frac{8\pi}{9}} Z_X^2 \alpha^2 \left(\frac{mc}{\hbar}\right)^2 \sqrt{\frac{E_0}{2mc^2}} |\bar{Q}_0|^2 F(Z_X Z, E_0) \rho_1. \quad (B12)$$

Here $g_s$ is the statistical weight factor depending on the angular momenta of the initial and final states, $m$ is the mass of the nucleus $N$ produced as a result of the fusion reaction and $E_0$ is its kinetic energy, which to a good accuracy coincides with the $Q$-value of the reaction. The factor $F(Z_X Z, E_0)$, defined as

$$F(Z_X Z, E_0) = \frac{|\psi_f(0)|^2_{Z=0}}{|\psi_f(0)|^2}, \quad (B13)$$

takes into account the deviation of the wave function of the final-state nucleus $N$ of charge $Z$ from the plane wave due to its interaction with the electric field of the X-particle. It is similar to the Fermi function employed in the theory of nuclear $\beta$-decay. By $\rho_1$ we denoted the quantity $|\psi_i(0)|^2$; it will be discussed in more detail below.

We estimate the transition matrix elements of the charge radius operator as

$$\bar{Q}_0 \simeq r_{i}, r_{f}. \quad (B14)$$

where $r_f \equiv \langle \psi_f^2 \rangle_f^{1/2}$ is the rms charge radius of the final-state nucleus, and $r_i$ is the rms charge radius of the compound nucleus in the initial state, which we express through the rms charge radii $r_{N_1}$ and $r_{N_2}$ of the fusing nuclei $N_1$ and $N_2$ according to eq. (A13). Thus, we actually estimate the transition matrix element of the charge radius operator $\bar{Q}_0$ as the geometric mean of the charge radii of the initial and final nuclear states.

The function $F(Z_X Z, E_0)$ can be written as

$$F(Z_X Z, E_0) = \frac{2\pi\eta}{1 - e^{-2\pi\eta}}. \quad (B15)$$

The Sommerfeld parameter $\eta$ is in this case

$$\eta = Z_X Z \alpha \sqrt{\frac{mc^2}{2E_0}}. \quad (B16)$$
Table II. Binding energies $E_b$, relative kinetic energies $E$ of fusing nuclei, astrophysical S-factors $S(E)$, reaction factors $A(E)$, $\rho$-parameters and rates $\lambda$ for the XCF reactions of stages I, II and III. Rates are inclusive of all sub-channels with either free or bound X-particles in the final state, except for IC processes, where final-state $X$ can only be free. $\rho$-factors shown in the sixth column are values of $\rho_1$ for IC reactions (3c) and (6c) and $\rho_0$ for all other reactions. See text for details.

| Reaction | $E_b$ (MeV) | $E$ (MeV) | $S(E)$ (MeV b) | $A(E)$ (cm$^3$/s) | $\rho_{0,1}$ (10$^{35}$ cm$^{-3}$) | $\lambda$ (s$^{-1}$) |
|----------|-------------|------------|-----------------|-------------------|-----------------|-----------------|
| $(ddX) \rightarrow ^4\text{He} + n + X$ | 0.290       | 0.145      | 0.102 [78]      | 1.31 \cdot 10^{-16} | 0.64          | 8.4 \cdot 10^{18} |
| $(ddX) \rightarrow ^3\text{He} + p + X$ | ”           | ”          | 8.6 \cdot 10^{-2} [78] | 1.11 \cdot 10^{-16} | ”             | 7.1 \cdot 10^{18} |
| $(ddX) \rightarrow ^4\text{He} + \gamma + X$ | ”           | ”          | 7 \cdot 10^{-9} [81] | 9.0 \cdot 10^{-24} | ”             | 5.8 \cdot 10^{11} |
| $(ddX) \rightarrow ^4\text{He} + X$ | ”           | ”          | 1.9 \cdot 10^{-2} | 2.5 \cdot 10^{-17} | 0.73          | 1.8 \cdot 10^{18} |
| $(^3\text{He}dX) \rightarrow ^4\text{He} + p + X$ | 0.91        | 0.36       | 7.1 [82]        | 4.0 \cdot 10^{-15} | 7.6 \cdot 10^{-3} | 3.0 \cdot 10^{18} |
| $(^3\text{He}dX) \rightarrow ^4\text{He} + n + X$ | 0.32        | 0.13       | 5.6 [83]        | 6.2 \cdot 10^{-15} | 0.88          | 5.5 \cdot 10^{20} |
| $(^3\text{He}dX) \rightarrow ^6\text{Li} + \gamma + X$ | 1.16        | 0.39       | 1.3 \cdot 10^{-8} [80] | 6.6 \cdot 10^{-24} | 1.8 \times 10^{-2} | 1.2 \cdot 10^{10} |
| $(^3\text{He}dX) \rightarrow ^6\text{Li} + X$ | ”           | ”          | 1.55           | 7.9 \cdot 10^{-16} | 0.14          | 1.1 \cdot 10^{19} |

Note the similarity of eq. (B15) with eq. (B3); the difference is that eq. (B3) describes the Coulomb repulsion of the like-sign charged nuclei $N_1$ and $N_2$, whereas eq. (B15) accounts for the Coulomb attraction of the oppositely charged X-particle and final-state nucleus $N$ of charge $Z$. This attraction increases the value of the atomic wave function of $N$ at $r = 0$ and leads to the enhancement of the reaction probability (Sommerfeld enhancement [86]).

The quantity $\rho_1$ in eq. (B12) is the squared modulus of the wave function of the compound nucleus in the initial state of the IC reaction, taken at zero separation between the nucleus and the X-particle. We evaluate it by integrating the squared wave function of the initial atomic state $(N_1N_2X)$ over the distance $\vec{r}$ between the fusing nuclei in the volume $|\vec{r}| \leq R_{1c} + R_{2c}$ (where $R_{1c}$ and $R_{2c}$ are the radii of the fusing nuclei) and setting the distance $R$ between their center of mass and the X-particle to zero:

$$\rho_1 \equiv \int_{|\vec{r}| \leq r_0} |\psi_1(\vec{R} = 0, \vec{r})|^2 d^3r, \quad r_0 = R_{1c} + R_{2c}. \quad (B17)$$

For our estimates we use the wave function $\Psi_1(\vec{r}_1, \vec{r}_2)$ defined in eq. (B6). Going from the coordinates $\vec{r}_1$ and $\vec{r}_2$ of $N_1$ and $N_2$ to their relative coordinate $\vec{r}$ and c.m. coordinate $\vec{R}$ and substituting into (B17), we obtain

$$a_s \equiv \left\{ \frac{m_2}{m_1 + m_2} a_1^{-1} + \frac{m_1}{m_1 + m_2} a_2^{-1} \right\}^{-1}. \quad (B18)$$

**Reaction** $(ddX) \rightarrow ^4\text{He} + X$ (3c). For this reaction $Z_1 = Z_2 = 1$, $Z = 2$, and $Q = 23.56$ MeV. The transition matrix element of the charge radius operator, estimated according to eqs. (B14) and (A13), is $Q_0 \simeq 4.52$ fm$^2$. For transitions from the $(ddX)$ state one has to take into account that the initial state of two spin-1 deuterons in the atomic $s$-state can have total spin $S = 2$ or $0$ (spin 1 is excluded by Bose statistics). This gives 6 possible initial spin states. As the final-state nucleus $^4\text{He}$ has zero spin and the transition operator is spin-independent, the IC transition (3c) is only possible from the $S = 0$ state of $(ddX)$. Therefore, $g_s = 1/6$. For evaluating the parameter $\rho_1$ given in eq. (B18) we use the values of $a_1 = a_2 = 8.53$ fm found from the variational treatment of $(ddX)$ atom with wave function (B6).

**Reaction** $(^4\text{He}dX) \rightarrow ^6\text{Li} + X$ (6c). In this case $Z_1 = 2$, $Z_2 = 1$, $Z = 3$, and $Q = 0.320$ MeV. For the transition matrix element of the charge radius operator we find $Q_0 \simeq 6.32$ fm$^2$. The reaction $(^4\text{He}dX) \rightarrow ^6\text{Li} + X$ is an E0 transition between nuclear states of total spin 1, therefore the weight factor $g_s = 1$. For our evaluation of $\rho_1$ we take $a_1 = 1.81$ fm, which is the Bohr radius of $(^4\text{He}X)$ atom, and $a_2 \simeq 30a_1$, as discussed in sec. A 1 c.

The expression for the rates of IC processes can conveniently be written in the form similar to (B5): $\lambda_{IC} = A \rho_1$, where the reaction factor $A$ is defined as the factor multiplying $\rho_1$ in eq. (B12). The values of the IC reaction factor $A$ and of the quantity $\rho_1$ for reactions (3c) and (6c) are presented in Table II, along with the reaction factors and rates of the other discussed XCF reactions.
To assess the accuracy of our calculations of the IC reaction factors, we compared our result for the \( (^{4}\text{He}X) \rightarrow ^{6}\text{Li} + X \) process with the existing calculations, which were carried out in the catalyzed BBN framework for the case of a singly charged catalyst particle \( C \) using a simple scaling law \[63\] and within a sophisticated coupled-channel nuclear physics approach \[87\]. To this end, we re-calculated our result taking \( Z_{X} = 1, \ Q = 1.3 \text{ MeV} \) and the c.m. energy \( E = 10 \) keV which were used in \[63, 87\]. For the reaction factor \( A \) of the process \( d + (^{4}\text{He}C) \rightarrow ^{6}\text{Li} + C \) we found \( A \equiv \lambda_{C}/\rho_{1} \approx 9.9 \times 10^{-17} \text{ cm}^{3}/\text{s} \). Eq. (B4) then gives for the corresponding astrophysical \( S \)-factor \( S(E) = 0.19 \) MeVb. This has to be compared with the results of ref. \[63\] (0.3 MeVb) and ref. \[87\] (0.043 MeVb). Our result lies between these two numbers, being a factor of 1.6 smaller than the former and a factor of 4.4 larger than the latter.

4. Branching ratios

The rates of the sub-channels of the XCF reactions in which the final-state \( X \) sticks to one of the produced nuclear fragments are obtained by multiplying the total rate of the channel, given in Table II, by the corresponding sticking probability \( \omega_{s} \), shown in Table III. The rate of the sub-channel with a free \( X \) in the final state is then found by subtracting from the total rate of the channel the rates of all the sub-channels with bound \( X \) in the final state. As an example, the rates of reactions (2b) and (2c) are obtained by multiplying the total rate of the \((ddX)\) fusion process with production of \(^{3}\text{H} \) and \( p \), \( \lambda = 7.1 \times 10^{18} \text{ s}^{-1} \), by \( \omega_{s} = 0.15 \) and \( \omega_{s} = 1.2 \times 10^{-2} \), respectively; the rate of reaction (2a) is then given by \( \lambda(1 - 0.15 - 1.2 \times 10^{-2}) \). It is then straightforward to find the branching ratios of all the discussed reactions; the results are presented in Table III.
| Reaction | $Q_0$ (MeV) | $Q$ (MeV) | $\omega_n$ | $\omega_s$ | $Br_0$ | $Br$ |
|----------|-------------|-----------|------------|------------|--------|------|
| $(ddX) \rightarrow ^3He + n + X$ | 3.27 | 2.98 | - | - | 54.2% | 29.1% |
| $(ddX) \rightarrow (^3HeX) + n$ | - | 3.89 | 0.61 | 0.40 | - | 19.4% |
| $(ddX) \rightarrow ^3H + p + X$ | 4.03 | 3.74 | - | - | 45.8% | 34.4% |
| $(ddX) \rightarrow (^3HX) + p$ | - | 4.01 | 0.22 | 0.15 | - | 6.2% |
| $(ddX) \rightarrow ^3H + (pX)$ | - | 3.84 | $1.9 \cdot 10^{-2}$ | $1.2 \cdot 10^{-2}$ | - | 0.5% |
| $(ddX) \rightarrow ^4He + \gamma + X$ | 23.85 | 23.56 | - | - | $3.7 \cdot 10^{-8}$ | $4 \cdot 10^{-9}$ |
| $(ddX) \rightarrow (^4HeX) + \gamma$ | - | 24.71 | 0.95 | 0.87 | - | $3 \cdot 10^{-8}$ |
| $(ddX) \rightarrow ^4He + X$ | - | 23.56 | - | - | - | 10.4% |
| $(^3He dX) \rightarrow ^4He + p + X$ | 18.35 | 17.44 | - | - | 100% | 94% |
| $(^3He dX) \rightarrow (^4HeX) + p$ | - | 18.60 | 0.29 | 0.06 | - | 6% |
| $(^3He dX) \rightarrow ^4He + (pX)$ | - | 17.54 | $2.3 \cdot 10^{-3}$ | $3.0 \cdot 10^{-4}$ | - | $3.0 \cdot 10^{-4}$ |
| $(^3HdX) \rightarrow ^4He + n + X$ | 17.59 | 17.27 | - | - | 100% | 96% |
| $(^3HdX) \rightarrow (^4HeX) + n$ | - | 18.42 | 0.23 | $4.0 \cdot 10^{-2}$ | - | 4.0% |
| $(^4He dX) \rightarrow ^6Li + \gamma + X$ | 1.475 | 0.32 | - | - | 100% | $10^{-13}$ |
| $(^4He dX) \rightarrow (^6LiX) + \gamma$ | - | 2.39 | $1 - 1.2 \cdot 10^{-6}$ | $1 - 1.2 \cdot 10^{-3}$ | - | $1.9 \cdot 10^{-8}$ |
| $(^4He dX) \rightarrow ^6Li + X$ | - | 0.32 | - | - | - | $\approx 100%$ |

Table III. General characteristics of reactions (1a)-(6c). $Q$ and $Br$ are $Q$-values and branching ratios of XCF reactions, $Q_0$ and $Br_0$ are values of these parameters for the corresponding X-less processes. $\omega_n$ and $\omega_s$ are X-sticking probabilities found for nuclei of finite and zero radius, respectively.
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