Chemical effects induced by the mechanical processing of granite powder

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Starting from 1970s, the use of mechanical forces to induce chemical transformations has radically changed vast areas of metallurgy and materials science. More recently, mechanochemistry has expanded to core sectors of chemistry, showing the promise to deeply innovate chemical industry while enhancing its sustainability and competitiveness. We are still far, however, from unveiling the full potential of mechanical activation. This study marks a step forward in this direction focusing on the chemical effects induced on the surrounding gaseous phase by the mechanical processing of granite. We show that fracturing granite blocks in oxygen can result in the generation of ozone. The refinement of coarse granite particles and the friction between fine ones are also effective in this regard. Combining experimental evidence related to the crushing of large granite samples by uniaxial compression and the ball milling of coarse and fine granite powders, we develop a model that relates mechanochemical ozone generation to the surface area effectively affected by fracture and frictional events taking place during individual impacts. We also extend the investigation to gaseous phases involving methane, oxygen, benzene and water, revealing that chemical transformations occur as well.

Mechanochemistry can be broadly defined as the branch of Chemistry that studies the chemistry activated and driven by the application of mechanical forces to chemical systems including, at least, one solid phase that is deformed beyond its yield point1–5. At present, this field of study is experiencing a period of remarkable flourishing characterized by the exponential growth of dedicated literature and the undertaking of international actions aimed at coordinating research and innovation activities6,7.

The unique mix of fundamental questions and practical applications that characterizes the mechanical activation of solids has attracted enormous interest starting from 1970s, when the first generation of oxide-dispersion-strengthened superalloys were fabricated8–11. A second, most marked surge of interest occurred in the 1990s, when the preparation of amorphous alloys and nanostructured materials by ball milling (BM) radically changed powder metallurgy and materials science in general12–21.

In the recent years, the mechanical processing of powders by BM has gradually expanded to inorganic and organic synthesis, thus approaching the broadest areas of chemistry. Due to the evident promise of bringing innovation in the fine chemical industry and enhancing its sustainability, mechanochemistry has been included among the ten technological innovations that can change our world22. To this aim, however, a full characterization of mechanochemical reactors and processing conditions, still fragmentary, is needed23–30, and a deeper understanding of the chemical reactivity induced by mechanical forces must be achieved. In this regard, kinetic investigations aimed at relating microscopic processes underlying mechanochemical transformations to processing variables and conditions are key to success31–47.

In general, the application of a mechanical force to a solid generates mechanical stresses that, in turn, give rise to mechanical deformation. As the solid is deformed beyond its elastic limit, chemical bonds get involved in deformation and the breakage of some of them can help accommodating local strain. Breaking and reforming of chemical bonds in sequence allows the nucleation and propagation of dislocations, and dislocation slip mediates plastic deformation48.

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Continued deformation generates new dislocations and the increase of dislocation density makes the solid increasingly harder due to dislocation interference. Progressively, the solid develops such resistance to further plastic deformation that the yield stress tends to become larger than the fracture stress. Under such conditions, cracks nucleate and grow. Eventually, the solid undergoes fracture and breaks apart in two or more fragments.

The rupture of chemical bonds during dislocation motion, crack generation and growth, and fracture clearly highlights the robust bridge connecting mechanics with chemistry. However, there is much more in this respect. Together with local thermal effects, the opening of new surfaces on short time scales in ionic solids can excite electromagnetic phenomena involving the emission of neutral and charged particles and photons. Charge separation due to fracture can generate very intense electric fields and electrons can be ejected from surface by field emission, which can also result in luminescence and electromagnetic waves. Tribological events can give rise to similar behaviours. Even at low stresses and sliding speeds, frictional electrification can produce enough electric charges to induce the dielectric breakdown of gases with consequent electric discharges and photon emission. The phenomena mentioned above can have immediate consequences for gaseous chemical species surrounding the deformed solid. In particular, they can induce chemical reactions in the gas phase as a result of the direct interaction between the excited solid surface and the molecules around. This makes the chemical transformations in gaseous phase activated by mechanical deformation a natural subject for mechanochemistry.

In this work, we provide quantitative evidence of the mechanochemical effects caused by the fragmentation and frictional processes undergone by a simple mineral such as granite. To this aim, we use BM.

Much of the undeniable success of BM is due to its ease of operation. Once balls and powder have been placed inside the reactor, this is sealed under the desired atmosphere and clamped to the ball mill. Depending on the device, balls and powder are set in motion by the movement of the reactor itself or of its mobile parts. After a certain time interval, BM is stopped, the reactor opened and the powder inside sampled or retrieved.

Nevertheless, such apparent simplicity hides considerable complexity. Powder is vigorously stirred and its particle dynamics becomes more and more cooperative and dissipative as the powder amount increases. Stirring and blending are enhanced by the milling balls, which also collide with each other and with the reactor walls. The powder particles trapped between the colliding surfaces undergo non-confined dynamic compaction at relatively high strain rates. Mediated by force chains formed across the volume of trapped powder, localized severe deformation processes take place, rapidly leading to fracture in the case of brittle solids.

The processes mentioned above take place concurrently and involve a broad range of length and time scales. In spite of this, we show that detailed information on the fragmentation of brittle materials and on the chemistry activated by fragmentation and frictional events can be still obtained. In particular, we use BM experiments to evaluate the amount of coarse granite powder that can be affected by fracture during individual impacts and the related increase of specific surface area induced by fragmentation. Granite was chosen because of its effectiveness in generating chemical effects in the surrounding atmosphere. In particular, in agreement with literature, we observe that surfaces formed by fracture activate gaseous oxygen, determining the formation of ozone. We relate the degree of chemical conversion of O₂ to O₃ to individual impacts and provide a self-consistent kinetic description of the mechanochemical processes underlying O₃ generation. Eventually, we extend the investigation to gaseous methane and to equimolar CH₄–O₂ and benzene C₆H₆–water H₂O gaseous mixtures, showing that chemical transformations are triggered also in these cases.

**Preliminary experiments on granite blocks under O₃ atmosphere.** Previous work has shown that the fracture of igneous and metamorphic rocks in air can produce O₃ at ambient pressure. Up to 4500 ppbv of O₃ were instantaneously detected, and a total of 0.4 μmol were generated, from the fracture of a 30×30×15 mm³ granite block inside a 5-dm³ isolated chamber under atmospheric pressure conditions and with background O₃ around 1.5 ppbv.

We carried out similar fracture experiments with 30×30×30 mm³ granite cubes sealed in a plastic container about 216 cm³ in volume under pure O₃ gas at ambient pressure (see Supplementary Information for details). Granite blocks were subjected to uniaxial compression until fracture occurred. The atmosphere was immediately sampled and analyzed using an O₃ monitor device. Measured O₃ levels, $c_{O₃}(t)$, as a function of time, $t$, are shown in Fig. 1 as a function of time, $t$. It can be seen that O₃ concentration suddenly rises as fracture occurs and, then, decreases with an...
exponential-like trend, reasonably due to ambient processes involving recombination of O atoms with either the container walls or the granite surface. After about 500 s, O3 concentration approximately goes back to the initial background value.

Experimental datasets can be satisfactorily best-fitted assuming that a maximum O3 concentration, $c_{oz,max}$, is produced instantaneously and that it decays exponentially according to the equation

$$c_{oz}(t) = c_{oz,max} \exp (-k_{rec}t) + c_{oz,0},$$

where $c_{oz,0}$ is the background O3 level and $k_{rec}$ measures the O3 recombination rate. In agreement with experimental measurements, the best-fitted curve suggests that, on the average, the fracture of the granite block results in a total concentration, $c_{oz,max}$, around 2.6 mmol m$^{-3}$, which corresponds to a total of about 0.56 μmol of O3 generated. The background O3 level, $c_{oz,0}$, is around 1.8 ppbv, whereas the best-fitted $k_{rec}$ value is equal to 0.010 s$^{-1}$.

We also evaluated the specific surface area of the granite samples before and after fragmentation (see Supplementary Information for details). The difference accounts for the surface area opened by the fracture process and is equal to about 169 cm$^2$. Therefore, the amount of O3 generated per unit surface area generated by fracture is around 33.1 μmol m$^{-2}$.

**Mechanical processing of granite powder under inert atmosphere.** Preliminary BM experiments have been performed to monitor the gradual refinement of powder particles caused by repeated impacts. To this aim, large granite pieces were crushed manually into shards around 2 cm in size. In a first set of experiments, shards were subjected to BM for very short time intervals in order to obtain coarse granules, which were subsequently separated from finer fractions. In a second set, BM was prolonged to refine the particle size to the finest possible fraction (see Supplementary Information for details).

In all cases, BM experiments were carried out using 5 g of granite powder and a cylindrical hardened steel reactor 2.0 cm in diameter containing a single 20-g stainless steel ball approximately 1.8 cm in diameter. This means that the ball adapts almost perfectly to the 1.0-cm curvature radius of the hemispherical bottom base of the reactor (see Supplementary Information for details), which allows the powder to be effectively trapped at impact and processed. Once sealed under inert Ar atmosphere, the reactor was subjected to a vertical harmonic oscillation with amplitude and frequency suitably set to make the ball collide with the bottom of the reactor every 2 s with an impact velocity of about 2.83 m s$^{-1}$ (see Supplementary Information for details).

The particle size of coarser and finer powders was measured by scanning electron microscopy (SEM). Representative SEM micrographs are reported in Fig. 2a,b. It can be seen that coarse powder exhibits average particle...
size around 1 mm and a relatively broad size distribution, whereas the powder subjected to prolonged BM has much finer particle size and narrower size distribution (see Supporting Information for details).

SEM observation carried out on powder sampled at intermediate time intervals during long BM experiments reveals that particle size refinement takes place gradually. The process was monitored performing systematic measurements of the granite powder surface area according to the Brunauer–Emmett–Teller (BET) method (see Supporting Information for details). It can be seen from Fig. 2c that the total surface area increases progressively with the number of impacts, \( m \). The observed behaviour can be satisfactorily described by the equation (see Supporting Information for details)

\[
S_{gr}(m) = (1 - k)^m S_{gr, in} + \left[ 1 - (1 - k)^m \right] S_{gr, fin}
\]

where \( S_{gr, in} \) and \( S_{gr, fin} \) are the initial and final values of surface area and \( k \) is the apparent rate constant of surface area generation. According to best-fitting, the quantities \( S_{gr, in} \) and \( S_{gr, fin} \) have values equal to 1.10 ± 0.02 m\(^2\) and 20.57 ± 0.01 m\(^2\) respectively. The apparent rate constant, \( k \), is equal to 2.1 × 10\(^{-2}\) ± 0.3 × 10\(^{-2}\). It follows that the first impact induces an increase in total surface area of about 0.41 m\(^2\).

The gaseous atmosphere was also sampled periodically during the mechanical processing using a gastight syringe. Through a gastight septum connector, the sampled gas was injected into a small chamber directly connected with the O\(_3\) monitor device. In agreement with expectations, no O\(_3\) was detected.

**Mechanical processing of coarse granite powder under O\(_2\) atmosphere.** A first set of BM experiments was carried out using 5 g of coarser granite granules and the 20-g ball. The reactor was sealed under Ar atmosphere and, subsequently, Ar was replaced with O\(_2\) (see Supporting Information for details).

Initially, experiments consisted of single impacts. After the impact, the gaseous atmosphere was analyzed every 10 s in order to evaluate the amount and lifetime of O\(_3\) produced. Results are shown in Fig. 3a. The O\(_3\) monitor device indicates that O\(_3\) concentration increases steeply immediately after the impact, reaches a maximum and, then, decreases to the background level with an exponential-like trend. After 500 s, the O\(_3\) concentration decay is complete. Overall, the observed behaviour is similar to the one of O\(_3\) generated by fragmentation of large

**Figure 3.** (a) Measured O\(_3\) concentration, \( c_{O_3}(t) \), as a function of time, \( t \). Data refer to experiments in which coarse granite powder is subjected to single impacts. (b) Measured O\(_3\) concentration, \( c_{O_3}(t) \), as a function of the number of impacts, \( m \). Data refer to experiments in which coarse granite powder is subjected to prolonged mechanical processing. (c) Measured O\(_3\) concentration, \( c_{O_3}(t) \), as a function of time, \( t \). Data refer to experiments in which fine granite powder is subjected to prolonged mechanical processing in the absence of milling ball. Best-fitted curves are shown. In (b), the red curve does not account for the O\(_3\) generation by frictional processes involving powder particles, whereas the green curve takes into account the frictional contribution.
granite blocks. Equation 1 can be used profitably to best fit the experimental data. The maximum O3 concentration, \(c_{oz,max}\), is around 48.1 mmol m\(^{-3}\) and the background O3 level, \(c_{oz,0}\), is approximately equal to 4.5 ppbv. The best-fitted \(k_{rec}\) value is found equal to 0.008 s\(^{-1}\).

Taking into account that the reactor has a volume of about 264 cm\(^3\), the total amount of O3 produced as a result of a single impact averages around 12.7 μmol. According to the experiments performed under Ar atmosphere, a single impact on coarse granite granules generates approximately 0.41 m\(^3\) of fresh surface area. Therefore, the amount of O3 produced per unit surface area generated by fracture during the impact is equal to about 30.9 μmol m\(^{-2}\). This value is quite close to the one of 33.1 μmol m\(^{-2}\) obtained from fracture experiments of granite blocks under O2 atmosphere.

We can reasonably deduce that the amount of O3 generated by fracture per unit surface area does not vary significantly with experimental conditions. This means that the amount of O3 detected after single impacts can be utilized to estimate the surface area generated by fracture and vice versa. For simplicity, we assume that the amount of O3 generated per unit surface area is equal to 33.1 μmol m\(^{-2}\). Hereafter, we denote it by \(k_f\).

Prolonged BM experiments involving coarse granite powder were performed to monitor the O3 production by fracture on the long term. Atmosphere was sampled every 10 s and analyzed with the O3 monitor device. The measured O3 concentration is plotted in Fig. 3b as a function of the number of impacts, \(m\). It can be seen that it increases smoothly, then reaches a maximum and, finally, attains a plateau value of about 15.5 mmol m\(^{-3}\).

In principle, the observed variation of O3 concentration can be explained taking into account three different contributions, namely the O3 generated by fracture, the O3 recombined at reactor walls or on the granite surface and the background O3 level. The three contributions can be readily included in a simplified modeling description of the O3 concentration variation that takes into due account the discrete nature of BM and, then, of O3 production (see Supplementary Information for details). Specifically, the O3 generated in a given impact is assumed proportional to the change in surface area caused by fracture and the decay process of the O3 amount generated is assumed to start immediately after the impact. The result is the expression

\[
c_{oz,f}(m) = k_{f} k (S_{gr,fin} - S_{gr,in}) \exp \left[-k_{rec}(m-1)\frac{1}{1-(1-k)\exp(-k_{rec}\tau)}\right] + c_{oz,0} \tag{3}
\]

where \(k_{rec}\) is the rate constant for O3 recombination and \(\tau\) is the time interval separating two consecutive impacts.

Equation 3 relates the O3 generation to individual impacts. In principle, it can be used to predict the variation of \(c_{oz}\) as the number of impacts increases. The time interval \(\tau\) between consecutive impacts is known and \(k_f, S_{gr,in}, S_{gr,fin}, k_{rec}\) and \(c_{oz,0}\) have been evaluated in previous experiments. However, we cannot exclude that the rate of recombination process, \(k_{rec}\), changes in the presence of repeated impacts and powder stirring. For this reason, we use Eq. (3) to best-fit the experimental points leaving \(k_{rec}\) free to vary as the only best-fitting parameter.

As shown in Fig. 3b, Eq. (3) is able to capture the general trend of experimental data. With a best-fitted \(k_{rec}\) value of about 0.014 s\(^{-1}\), it well reproduces the initial increase, the maximum and the subsequent exponential decay. However, it is definitely unable to match the final plateau value.

In this respect, it is worth noting that the final values attained by O3 concentration are much higher than the expected background level, \(c_{oz,0}\), much lower than 15.5 mmol m\(^{-3}\). The solution to this apparent inconsistency comes from the experiments performed on finer granite powder.

Experiments were carried out, both in the presence and in the absence of the milling ball, under O2 atmosphere for long mechanical processing times. The reactor atmosphere was analyzed every 10 s with the O3 monitor device. The O3 concentration measured during experiments carried out in the absence of milling ball is shown in Fig. 3c as a function of time. Experimental data reveal that O3 concentration increases smoothly, reaching a plateau value of about 15.3 mmol m\(^{-3}\).

This value is significantly higher than the background O3 level observed in experiments involving single impacts and substantially coincident with the final plateau value observed in experiments involving the prolonged mechanical treatment of coarser granite powder. Approximately the same behaviour is observed both in the presence and in the absence of the milling ball. We infer that O3 generation takes place despite no significant fracture process occurs to modify the particle size of granite powder. Therefore, we ascribe the observed O3 generation to frictional processes involving the surface of powder particles.

Based on such hypothesis, a suitable modeling description can be developed (see Supporting Information for details). Experimental data can be satisfactorily best-fitted by the expression

\[
c_{oz,\alpha}(m) = k_{\alpha} \exp \left[-k_{rec}(m-1)\right] \left[ S_{gr,fin} \frac{1}{1-(1-k)\exp(-k_{rec}\tau)} - (S_{gr,fin} - S_{gr,in})(1-k) \frac{1}{1-(1-k)\exp(-k_{rec}\tau)} \right] + c_{oz,0} \tag{4}
\]

which allows estimating the amount of O3 generated by attrition per unit surface area, \(k_{\alpha}\). The best-fitted \(k_{\alpha}\) estimate is equal to about 0.02 μmol m\(^{-2}\). Therefore, it is three orders of magnitude smaller than the amount of O3 generated by fracture surfaces, \(k_f\), which has been found equal to about 33.1 μmol m\(^{-2}\).

If we assume that friction between particles generates O3 also during the mechanical processing of coarse granite powder, Eq. (3) has to be re-written as (see Supplementary Information for details)

\[
c_{oz}(m) = c_{oz,f}(m) + c_{oz,\alpha}(m). \tag{5}
\]

In fact, Eq. (5) best fits the experimental data in Fig. 3b with a background O3 level, \(c_{oz,0}\), of about 5.3 ppbv. The estimated values of the different quantities are summarized in Table 1. It can be seen that there is substantial agreement between the values obtained in different experiments. It follows that systematic experiments
The amount of $O_3$ generated by fracture per unit surface area, $\rho_{O_3}$, is approximately equal to $1.4 \times 10^{10}$ cm$^{-2}$ and $2.9 \times 10^{10}$ cm$^{-2}$ respectively. These values can be compared with the surface density of active sites of about $1.9 \times 10^{19}$ cm$^{-2}$. This value is definitely higher than the surface density of ions in surfaces freshly generated by fracture, which can be as high as $1.0 \times 10^{15}$ cm$^{-2}$.

Mechanical processing of granite powder under CH$_4$ and equimolar CH$_4$–O$_2$ atmospheres. If electromagnetic phenomena are involved in $O_3$ generation, the formation of new surfaces by fracture or friction between powder particles can be expected to drive to reaction chemical species different from $O_2$. To investigate this issue, we focused on the chemical response of gaseous CH$_4$. This choice is related to the availability of literature on the reactivity of gaseous CH$_4$, and organic compounds in general, in the presence of glow discharges$^{54,56}$. Experiments were performed under the same processing conditions utilized for experiments under $O_2$ atmosphere. Specifically, about 5 g of coarse granite granules were introduced in the reactor together with the 20-g ball. Then, the reactor was sealed under Ar atmosphere and Ar suitably replaced with CH$_4$. The gaseous atmosphere was sampled and analyzed by gas-chromatography using a flame ionization detector (see Supporting Information for details).

Experimental data shown in Fig. 4a clearly reveal the partial conversion of CH$_4$ to ethane (CH$_3$CH$_3$) and other light hydrocarbons. Therefore, the prolonged mechanical processing of granite by BM activates CH$_4$ molecules and allows their combination in more complex molecular units.

No kinetic modeling based exclusively on mechanochemical processes occurring during individual impacts can be developed in the present case. Indeed, the mechanochemical kinetics responsible for the generation of the chemical species that eventually allow the formation of hydrocarbons is inextricably intertwined with chemical kinetics in the gaseous phase. Therefore, no direct relationship can be expected between the concentration of gaseous species and the fragmentation and frictional processes undergone by granite.

Similar considerations hold for the experiments performed using an equimolar mixture of CH$_4$ and O$_2$ gases. In this case, gas-chromatographic analyses allow detecting CH$_2$CH$_3$ and light hydrocarbons as well as oxidized chemical species such as methanol (CH$_3$OH), dimethyl ether (CH$_3$OCH$_3$) and formaldehyde (CH$_2$O). The concentrations of a few chemical species are shown in Fig. 4b as a function of time. Chemical species appear, more or less, in order of increasing complexity, suggesting that simpler compounds combine with each other to form more complex ones. The concentration of the different chemical species increases as time increases.

Mechanical processing of granite powder under equimolar C$_6$H$_6$–H$_2$O atmosphere. In the attempt to further explore the chemistry of gaseous phases initiated by the mechanical processing of granite, we carried out prolonged BM experiments on granite powder in the presence of gaseous C$_6$H$_6$ and H$_2$O. To this aim, experiments were performed at 150 °C using the same processing conditions utilized for other atmospheres (see Supporting Information for details).

As shown in Fig. 4c, we observe the conversion of C$_6$H$_6$ to different complex species. Those with higher concentration are biphenyl (C$_{12}$H$_{10}$), phenol (C$_6$H$_5$OH) and 1,4-cyclohexadiene (C$_6$H$_8$). Again, the prolonged mechanical processing of granite shows the capability of inducing chemical reactivity in organic molecules.

Discussion

The mechanical processing of granite powder gives rise to chemical effects in the presence of reactive gases. Literature suggests that such effects can be ascribed to the electrons emitted as a consequence of the intense electric fields that are generated when a fracture separates electrical charges located on the parting surfaces$^{54}$. The heterolytic rupture of chemical bonds, and the consequent formation of ions, can result in electric fields of about $1.8 \times 10^6$ ρ V cm$^{-1}$, where ρ is the surface density of separating charges. The electric breakdown of air takes place at electric fields around 30 kV cm$^{-1}$, which require ρ values approximately equal to $1.7 \times 10^{10}$ cm$^{-2}$. The surface density of ions needed to activate discharges in gaseous O$_2$ and CH$_4$ is on the same order of magnitude, i.e. about $1.8 \times 10^{-6}$ μmol m$^{-2}$. Under the hypothesis that the number of O$_3$ molecules generated equals the number of active sites responsible for electromagnetic phenomena, such estimate indicates that we can expect a surface density of active sites of about $1.9 \times 10^{15}$ cm$^{-2}$. This value is definitely higher than the surface density of separating charges, ρ. The amount of O$_3$ generated by friction per unit surface area, ρ, is approximately equal to $33.1 \mu$mol m$^{-2}$.
to 0.02 μmol m⁻² and leads to a lower surface density of active sites, around 1.2 × 10¹⁶ cm⁻², which is however higher than ρ. It follows that both fracture and frictional processes can give rise to electric fields with intensity well beyond the threshold required to activate plasma discharges in the investigated gaseous phases. This means that electromagnetic phenomena activated by fracture and friction can explain, in principle, the observed O₃ generation and the chemical reactivity in the presence of CH₄ as well as of CH₄—O₂ and C₆H₆—H₂O equimolar mixtures.

In this regard, however, at least another mechanism can be invoked. Mechanical processing by BM has been shown to generate excited chemical bonds at the surface of solids that, in turn, determine the consumption of a radical scavenger. Specifically, the fracture and friction of quartz particles dispersed in liquid ethanol has been shown to generate active sites with surface density respectively equal to about 4.8 × 10¹³ and 2.8 × 10¹² cm⁻². The surface density of active sites formed by fracture is comparable, and possibly even higher, than the maximum possible surface density of dangling bonds, which is equal to about 1.2 × 10¹³ cm⁻² for quartz. This suggested a very high reactivity for quartz surfaces excited by mechanical activation, eventually resulting in the formation of radicals. Similar results have been obtained for pozzolanas in the presence of H₂O.

The formation of radical species can explain the reactions involving CH₄. In particular, we can hypothesize the formation of CH₃ radicals as a consequence of interaction of CH₄ molecules with the excited granite surface. We can also expect the formation of O radicals in the presence of O₂ molecules. Therefore, the organic molecules formed in the presence of CH₄ and O₂ can be the result of radical reactions.

In contrast, the formation of C₆H₈ in the presence of C₆H₆ and H₂O can be hardly explained in terms of a radical transformation. Rather, it is reminiscent of the old-fashioned Birch reduction, a widely used reaction in synthetic organic chemistry that converts benzenoid compounds into 1,4-cyclohexadienes. Although reaction conditions are quite different from conventional ones using ammonia and alkaline metals, we can still conjecture that the initial step involves the reduction of the aromatic ring by an electron. We simply remark that, in our case, electrons can be made available by electrification phenomena caused by fracture or attrition.
Conclusions

Experimental findings prove that the mechanical processing of granite can have significant chemical effects on the surrounding gaseous phase. In the presence of oxygen, fracturing and frictional processes determine the formation of relatively small amounts of ozone. The direct relationship between ozone generation and mechanical activation allows developing an accurate kinetic model that connects the kinetic evidence on the macroscopic scale with the number of impacts, the total surface area involved in fracturing and frictional events, and the characteristic ozone lifetime under the processing conditions investigated. The kinetic model bridges successfully the global and local scales of the mechanochemical transformation, showing that experimental kinetics can be exploited to obtain information on the local chemistry underlying mechanically activated reactions. It is precisely such local chemistry that most genuinely points to the heart of the chemistry induced by the application of mechanical forces.

The kinetic model can be extended to other mechanochemical transformations characterized by a direct relationship between mechanical activation and chemical effect. This is not the case of the mechanochemical reactions induced in the presence of gases more complex than oxygen. Indeed, the global kinetics also depends, in such cases, on the intrinsic gaseous phase behaviour. And yet, it is worth noting that the mechanical processing of a simple, apparently inert mineral, and presumably of other minerals and ionic solids, can give rise to significant chemistry. We ascribe the observed mechanically induced reactivity to the generation of intense electromagnetic fields. Seemingly, this is the most reasonable working hypothesis. However, we cannot hide the fact that investigation in this area is just beginning and we have yet to face the most important challenges.

Data availability

Datasets generated and analysed during the current study cannot be shared at this time as they also forms part of an ongoing study. However, they can be available totally or in part from the corresponding author on reasonable request.

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Author contributions
FD. designed the research; A.L.S., M.C., G.P., S.G., A.P., and F.D. conducted the experiments; F.D., S.G., and A.P. wrote the paper; All authors discussed and reviewed the manuscript.

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The authors declare no competing interests.

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