High-Power Ultrasonic Synthesis and Magnetic-Field-Assisted Arrangement of Nanosized Crystallites of Cobalt-Containing Layered Double Hydroxides

Andrei N. Salak 1,*, Daniel E. L. Vieira 1, Irina M. Lukienko 2, Yuriy O. Shapovalov 2, Alexey V. Fedorchenko 2, Elena L. Fertman 2, Yuriii G. Pashkevich 2,3, Roman Yu. Babkin 3, Aleksandr D. Shilin 4, Vasili V. Rubanik 4, Mário G. S. Ferreira 1, Joaquim M. Vieira 1

1 Department of Materials and Ceramics Engineering, CICECO – Aveiro Institute of Materials, University of Aveiro, 3810-193 Aveiro, Portugal
2 B. Verkin Institute for Low Temperature Physics and Engineering of the National Academy of Sciences of Ukraine, 47 Nauky Ave., 61103 Kharkov, Ukraine
3 O. Galkin Donetsk Institute for Physics and Engineering of the National Academy of Sciences of Ukraine, 46 Nauky Ave., 03680 Kyiv, Ukraine
4 Institute of Technical Acoustics of National Academy of Sciences of Belarus, Lyudnikov Avenue, 13, 21009 Vitebsk, Belarus

* Correspondence: salak@ua.pt; Tel.: +351-234-247318

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Abstract: High-quality stoichiometric Co2Al–NO3 and Co2Al–CO3 layered double hydroxides (LDHs) have been obtained by precipitation followed by anion exchange, both high-power-sonication assisted. Application of high-power ultrasound has been demonstrated to result in a considerable acceleration of the crystallization process and the anion-exchange reaction. Two independent approaches were used to form bulk and 2-D samples of Co2Al–NO3 with the oriented crystallites, namely uniaxial pressing of deposits from sonicated LDH slurries and magnetic field-assisted arrangement of LDH crystallites precipitating on glass substrates. A convenient way of preparation of semi-transparent compacts with relatively big blocks of oriented crystallites have been demonstrated. Thin dense transparent films of highly-ordered crystallites of Co2Al–NO3 LDH have been produced and characterized.

Keywords: magnetic layered structure; oxygen octahedron; anisotropic crystallite; highly-ordered arrangement

1. Introduction

Layered double hydroxides (LDHs) of the M2+–M3+ type represent the most numerous group in the anion clays family [1]. Although some of them occur as natural minerals, most of the known LDHs are synthetic. Their framework is composed of parallel positively-charged layers of the double metal hydroxides, \([\text{M}^2+_{1-x}\text{M}^3+x(OH)_2]\)^+, where \(0.17 \leq x \leq 0.50\) [2]. The positive charge of the hydroxide layer is compensated by anions \((\text{A}^y)^-\) intercalated into the interlayer gallery and screened from each other by water molecules [2]. To denote the chemical composition of an LDH, a short form, namely \(\text{M}^{2+}_n\text{M}^{3+}_n\text{A}^y\), where \(n = (1-x)/x\) is the atomic ratio of the cations \(\text{M}^2+\) and \(\text{M}^3+\) in the layer, is commonly used. Owing to the complex electrostatic interaction between hydroxide layers and the interlayer species, LDHs can be intercalated with anions and anionic complexes, which are very different in nature, size, geometry and charge [2–9]. Moreover, the same anions in the interlayer of the same LDH can be arranged in several ways depending on the cations ratio [9] and on the relative amount of...
intercalated water [10] that results in different distances between the adjacent hydroxide layers (basal spacings).

Layered double hydroxides are widely used in catalysis; however, the general applications of LDHs are resulted from their unique anion-exchange ability [3]. In their capacity of exchangers and adsorbents, LDHs are extensively used as (nano)containers for delivery of drugs, genes and markers in vivo [11–14], active pigments in coatings for corrosion protection [15–17], agents in the systems for water purification [18–20] and in other areas. In such applications, the nature of cations M$^{2+}$ and M$^{3+}$ is disregarded (except for apparent aspects of low cost and ecological compatibility), since in respect of anion-exchange properties the M$^{2+}$/M$^{3+}$ ratio is only important since it entirely determines the layer charge per unit cell [21].

Functionalities of LDHs can be extended by means of a use of particular cations in the hydroxide layers. Luminescence properties of LDHs with M$^{3+}$ partly substituted by rare earth cation [22–24] have recently been reported. The layered double hydroxides, in which at least one of the metal cations is magnetic: Co$^{2+}$, Ni$^{2+}$, Mn$^{3+}$, Cr$^{3+}$ or Fe$^{3+}$, were studied [25–33]. Onset of spontaneous magnetization and a peak of magnetic susceptibility were observed in those LDHs at temperatures of a few K. The highest Tc of 26 K was reported for Ni$_4$Mn–CO$_3$ LDH [28]. Because of the very low Tc values, direct practical exploration of the magnetic transition in such materials is unlikely. Nevertheless, some features of such magnetic LDHs, in particular the tunability of their magnetism via the layer cation content, the cation ratio and the interlayer distance can be used in preparation of functional materials. Coronado and co-workers have published several successful examples of the application of layered double hydroxides in formation of magnetic hybrids [29–32].

Unusual magnetic behavior of LDHs containing Co$^{2+}$, namely Co$_3$Al–NO$_3$ and Co$^2$Al–NO$_3$ in the temperature range above liquid nitrogen has recently been reported. Specifically, between about 80 and 150 K, both the paramagnetic Curie temperature and the effective magnetic moment were found to vary in a non-monotonous way [34]. Based on a detailed consideration of crystal structure and modeling, Babkin et al. have shown that freezing of tilting fluctuations of O–H bonds are responsible for these anomalies [35]. Taking into account the statistical distribution of divalent and trivalent cations, and their effect on the distortions of the original structure, the temperature dependences of the magnetic susceptibility of Co$_3$Al–NO$_3$ LDHs ($n = 2,3,4$) were calculated. It was noted that cobalt-containing LDHs represent convenient models for experimental modeling of effects of frustrated magnetic states in solids.

Unfortunately, the maximum size of the LDH crystallites, which are flake-like with the typical diameter-to-thickness ratio of about 20:1, usually does not exceed 1–2 $\mu$m; in the case of Co$_3$Al layered hydroxides, it is generally 50–100 nm [34]. Nevertheless, Abellán et al. [32] and Carrasco et al. [33] have prepared Co$_3$Al LDHs with relatively big (micrometric-size) regularly shaped crystallites.

The main problem that makes difficult any manipulations with individual LDH crystallites is their strong tendency to form shapeless agglomerates. The research group of Evans and Duan reported a use of a 0.2 $\mu$m membrane to filter the Co$_3$Fe LDH suspension with crystallites of about 100 nm in diameter, which then were assembled in an external magnetic field [36]. As a result, LDH–porphyrin composite film has been formed with a promising potential in application as an electrochemical sensor for glucose [37].

Co$^{2+}$-containing LDHs are generally produced using either the conventional co-precipitation methods (with the total duration of about 48 h) [31,32,34] or the hydrothermal synthesis (at least 24 h at 100–130 °C long) [33] or a combination of both [36,37]. Layered hydroxides synthesized in such ways contain anions of the initial metal salts or with other anions, which are present or created in the reaction solution. In the majority of applications, LDHs must be intercalated with functional anions: bioactive substances, corrosion inhibitors etc. The intercalation is implemented by means of anion exchange in the respective solutions. Complete substitution of the parent anions by functional ones takes from few hours to several days and even weeks depending on the anion’s nature.

In this work, high-power sonication was applied at different stages of the Co$_3$Al–NO$_3$ synthesis and the nitrate-to-carbonate anion exchange processes in order to accelerate the LDH production. For comparison, the same LDH compositions were obtained using the conventional co-precipitation
procedure followed by anion exchange. Besides, we used the sonication treatment to de-agglomerate the obtained LDH crystallites and to facilitate their arrangement on a substrate.

The main objective of this work was to order and assemble crystallites of Co₂Al–NO₃ and Co₂Al–CO₃ LDHs with an assistance of an external magnetic field. The composition with the $n = 2$ cations ratio was chosen because the parent Co₂Al layered double hydroxides with this $n$ value (i) are the easiest to obtain reproducibly and (ii) allow simple exchanges for numerous inorganic and organic anions. We have obtained semi-transparent compacts and transparent thin films of the arranged LDH crystallites and characterized them with respect to their morphology and some optical properties.

2. Materials and Methods

Cobalt (II) nitrate hexahydrate ($\geq 98\%$), aluminum nitrate nonahydrate ($\geq 98.5\%$), sodium hydroxide ($\geq 98\%$), sodium nitrate ($\geq 99.5\%$), sodium carbonate ($>99.95\%$) were purchased from Sigma-Aldrich, while nitric acid (65%) was purchased from Merck. The chemicals were used as received without further purification. All the solutions were prepared in deaerated water. The co-precipitation procedure was carried out under a nitrogen atmosphere. Proportions of the reagents were chosen to meet the desired molar cation ratio (Co/Al = 2) without any excess.

To prepare Co₂Al–nitrate LDH using the conventional method, a solution containing 0.5 M Co(NO₃)₂∙6H₂O and 0.25 M Al(NO₃)₃∙9H₂O was instilled to a 1.5 M NaNO₃ solution under continuous stirring at room temperature. During the precipitation, pH of the mixture was controlled by addition of a 2 M NaOH solution to be 8. This process took about one hour.

The obtained suspension was kept in a water bath at about 100 °C for 4 h to complete the crystallization and then centrifuged at 10⁴ rpm for 90 s. The resulting dense slurry was washed with deionized water followed by centrifugation. This procedure was repeated three times to remove a residual NaNO₃.

Sonication-assisted experiments were performed using a VCX 1500 Sonics processor (max output power 1.5 kW at 20 kHz) equipped with a modified high-volume continuous flow cell.

High-power sonication was applied at three different stages of the LDH production, namely (1) at co-precipitation (synthesis) step, (2) at crystallization step and (3) at anion-exchange stage. A certain amount of the suspension/slurry obtained after each step was centrifuged, washed and dried as described above then used for the phase analysis.

In the first experiment, it was attempted to substitute the relatively long co-precipitation step by the sonication-assisted direct synthesis. The solutions of Co(NO₃)₂∙6H₂O, Al(NO₃)₃∙9H₂O, NaNO₃ and NaOH in the same amounts as those used for the conventional co-precipitation were mixed under a 1.5 kW ultrasonic agitation for 3, 5 or 10 min.

In the second experiment, the Co₂Al–NO₃ LDH slurry was sonicated at 1.5 kW for 2, 5 and 10 min just after the conventional co-precipitation step (without crystallization in a water bath).

The third experiment was conducted aiming at acceleration of the nitrate-to-carbonate anion-exchange reaction. The Co₂Al–NO₃ LDH slurry after the co-precipitation and the crystallization in a water bath was mixed with a 0.1 M NaCO₃ solution at room temperature. The pH value of the mixture was adjusted to be 8 using a 2 M NaOH solution. The obtained solution was sonicated at 1.5 kW for 7 min and then centrifuged at $10^4$ rpm for 90 s. The LDH slurries prepared using the aforementioned procedures were estimated to contain about 85% of water.

For the phase analysis and the crystal structure characterization, the final product was dried at 60 °C for 24 h. The X-ray diffraction (XRD) data of the obtained LDH powders were collected using a Rigaku D/MAX-B diffractometer equipped with a graphite monochromator to reduce contributions caused by fluorescence of cobalt in Cu Kα radiation.

Two types of the samples, namely bulk samples (pressed compacts of LDH powders) and thin films (layers of the arranged LDH crystallites) were prepared with assistance of an external magnetic field. The experimental procedure was the following. A slurry of Co₂Al–NO₃ or Co₂Al–CO₃ LDH produced using the conventional co-precipitation route followed by 4-min high-power sonication was dispersed in ethanol. The procedure was carried out in ultrasonic bath for 15 min. For the bulk samples preparation, the slurry-to-ethanol ratio was 0.05 g/mL, while the thin films were deposited
using the suspension with the ratio of 0.01 g/mL. Immediately after the ultrasonic treatment, the suspension was put in a flat 10 mm height plastic container with a 15 × 15 × 0.5 mm glass substrate at the bottom. The container was placed horizontally between the vertically oriented poles of a GMW Model 3470 electromagnet connected to an Agilent Model 6811B Power Source/Analyzer. The magnetic field generated in the pole gap was 0.5 T. The field was applied in three cycles of 1 min each. The cycles were divided into pulses of 10 s for the first minute and 20 s for the next two cycles. Then the substrate with the precipitated LDH was dried in the oven at 60 °C until the complete evaporation of ethanol. For comparison, the samples were also prepared using the same conditions in a zero magnetic field.

To prepare the bulk samples, thick layers of the deposited LDH particles were carefully removed from the glass substrate, placed on the die of a 6 mm cylindrical mold and pressed into semi-transparent slabs of about 0.30 mm thick. Some as-pressed samples were additionally polished to achieve better transparency.

Morphology of the deposited LDH layers was studied using a Hitachi S4100 scanning electron microscope (SEM) with an electron beam accelerating voltage of 25 kV. Size and morphology of the LDH particles/crystallites were characterized using a Hitachi HD-2700 scanning transmission electron microscope (STEM) with an electron beam accelerating voltage of 200 kV.

3. Results and Discussion

XRD patterns of the Co2Al–NO3 LDH samples synthesized from a mixture of the respective nitrates either using the conventional co-precipitation route or by means of ultrasonic agitation of the mixture are shown in Figure 1.

![Figure 1](image_url)  

**Figure 1.** X-ray diffraction (XRD) patterns of the Co2Al–NO3 layered double hydroxides (LDHs) synthesized using the conventional co-precipitation method (a) and by means of the ultrasonic treatment of the stoichiometric mixture of the respective metal nitrates at 1.5 kW for 3 min (b), 5 min (c), and 10 min (d) followed by 4-h thermal treatment in a water bath. The strongest peak of the residual sodium nitrate (from the parent solution) is indicated.

It is seen from Figure 1 that in all the cases when sonication-assisted direct synthesis from the mixture of the respective nitrates was attempted the Co2Al–NO3 LDH phase was not formed regardless of the sonication treatment time. Although the most intense diffraction reflections associated with LDH structure were seen in the XRD patterns (Figure 1b–d), these reflections broadened and were shifted with respect to those typical of Co2Al–NO3 (Figure 1a). When one prepares LDH using the conventional co-precipitation route, the synthesis goes rather slowly, taking
about 1–2 hours and requires constant control and correction of pH. In the case of the direct sonication-assisted synthesis attempted in this work, in spite of the fact that the pH value was adjusted to the desired one in the starting mixture, it was impossible to control pH in the course of the high-power ultrasound treatment. We suggest that the pH value unpredictably changed during the sonication and this could have led to deviation in the cations ratio in the resulting LDH material as well as to intercalation of hydroxide anions together with (or instead of) nitrate anions.

A water bath thermal treatment of the suspension obtained by the co-precipitation is an important stage of formation of an LDH. At this stage, the LDH crystallite size is equalized. We have attempted to substitute this 4-hour stage by a high-power sonication for several minutes. XRD patterns of the respective LDHs are shown in Figure 2.

Figure 2. XRD patterns of the Co2Al–NO3 LDHs synthesized using the conventional co-precipitation method and then either kept in a water bath at about 100 °C for 4 h (a) or sonicated at 1.5 kW for 2 min (b), 5 min (c), and 10 min (d).

One can see in Figure 2 that the XRD patterns of the samples obtained after the high-power sonication treatment of the LDH suspension are similar to the XRD pattern of the Co2Al–NO3 LDH prepared from the same suspension using a water bath treatment. The similarity of the patterns is especially evident for the sample sonicated for 5 min. Hence, a 4-h stage of the water bath treatment can be effectively substituted by a 5-min sonication stage. We suggest that a considerable reduction of the LDH crystallization is possible due to the acoustic cavitation caused by high-power sonication [38]. Acoustic cavitation is the formation, growth and collapse of bubbles in a liquid under the influence of an ultrasonic field [39]. The implosive collapse of the bubbles produces a localized heating area via shock wave formation (adiabatic compression) within the gas phase of the collapsing bubble.

The results obtained in our experiment on the sonication-assisted crystallization of LDHs are in agreement with those reported in previous works on the preparation of nanomaterials using the ultrasonic waves [40–42]: indeed, application of high-power ultrasound considerably accelerates chemical processes in aqueous solutions.

Figure 3 shows the XRD patterns of the Co2Al–NO3 LDH before and after the nitrate-to-carbonate exchange in the solution of Na2CO3 and NaOH under high-power sonication for 7 min. The XRD pattern of the material obtained as a result of such a treatment corresponded entirely of that of Co2Al–CO3. It should be stressed here that the standard NO3− → CO32− anion-exchange reaction in Co2Al–NO3 takes about seven days of continuous stirring at room temperature.
Figure 3. XRD patterns of the Co₂Al–NO₃ LDH prepared using the conventional co-precipitation method before (a), after the NO₃⁻ → CO₃²⁻ anion-exchange reaction at room temperature either without sonication for 7 min (b), 60 min (c), 24 h (d) and 7 days (f) or under a 1.5 kW sonication for 7 min (e).

Such a considerable reduction of the time required to complete the anion-exchange reaction makes an application of the high-power sonication very promising.

It is known that anion-exchange is rather a diffusion process. We suggest that the high-power ultrasonic treatment enhances significantly the “interlayer diffusion” owing to both the local heating and the mechanical impacts on the hydroxide layers.

Lattice parameters (in a hexagonal setting) and basal spacings calculated from the 2Theta positions of the diffraction reflections (003), (006) and (110) for all the LDH compositions prepared in this work are listed in Table 1.

| LDH composition                                        | a (Å)  | c (Å)  | d (Å)  |
|--------------------------------------------------------|--------|--------|--------|
| Co₂Al–NO₃ Conventional co-precipitation followed by     | 3.063  | 26.732 | 8.911  |
| 4-h crystallization in a water bath                     |        |        |        |
| Co₂Al–NO₃ (sonication-assisted crystallization)         |        |        |        |
| 2 min                                                 | 3.049  | 26.585 | 8.862  |
| 5 min                                                 | 3.071  | 26.579 | 8.860  |
| 10 min                                                | 3.054  | 26.513 | 8.838  |
| Co₂Al–CO₃ Standard anion-exchange (7 days)              | 3.072  | 22.878 | 7.626  |
| Sonication-assisted exchange (7 min)                    | 3.070  | 22.831 | 7.610  |

The relative errors in determination of the parameters a and c were 0.07% and 0.1%, respectively.

One can see that the values of lattice parameters of the same LDH compositions prepared using either the conventional routes or the sonication-assisted procedures were rather similar. However, in cases of the application of high-power sonication, the values of lattice parameters were regularly lower. This can be caused by some deviations from the desired cations ratio, Co/Al = 2. It should be noticed that the lattice parameters values for Co₂Al–CO₃ LDHs available from the literature are even
lower than those obtained in our study (Table 1): cf. $a = 3.066 \text{ Å}, c = 22.593 \text{ Å}$ [43] and $a = 3.068 \text{ Å}, c = 22.609 \text{ Å}$ [44].

An optical image (in nonpolarized light) of a pressed compact of $\text{Co}_2\text{Al–NO}_3$ layered double hydroxide powder precipitated in an external magnetic field and then dried without mixing is shown in Figure 4a. For comparison, the respective image of a compact of the same LDH composition prepared using the conventional route followed by centrifugation, drying and mixing with mortar and pestle is demonstrated in Figure 4b.

![Figure 4](image1.png)

**Figure 4.** Images of the $\text{Co}_2\text{Al–NO}_3$ samples in nonpolarized light: the pressed compact of the LDH powder precipitated in an external magnetic field and then dried without mixing (a) and the pressed compact of the LDH powder dried after centrifugation (b).

Relatively big (up to 0.5 mm long) individual blocks with the same contrast are clearly seen in the samples pressed from the LDH powder deposited in an external magnetic field. Similar blocks are also observed in the $\text{Co}_2\text{Al–NO}_3$ sample pressed from the powder dried without precipitation in the field; however, their average size is considerably smaller. The same difference between average dimensions of the blocks visible in nonpolarized light was found in the respective compacts of $\text{Co}_2\text{Al–CO}_3$ LDHs.

Figure 5 shows the polarized light images of the $\text{Co}_2\text{Al–CO}_3$ compacts prepared from the LDH crystallites precipitated with and without an external magnetic field. The samples under study were polished to achieve thickness of about 0.15 mm.

![Figure 5](image2.png)

**Figure 5.** Images of the $\text{Co}_2\text{Al–CO}_3$ samples in a polarized microscope: the pressed compact of the LDH powder precipitated in an external magnetic field and then dried without mixing (a) and the pressed compact of the LDH powder dried after centrifugation (b). Inset shows the magnified image of the zone with the iridescent colors.

Since the thickness of these two samples is almost the same, their images in crossed polarizers can be compared by contrast. A higher contrast between dark and light areas suggests more homogeneous crystallite blocks (Figure 5). It is seen that the blocks of the sample prepared using the magnetic field assisted approach are bigger and more homogeneous than those in the sample produced via the conventional route. Iridescent color of some blocks is a manifestation of birefringence.
Figure 6 shows images of the zone with the iridescent colors observed in the Co$_2$Al–CO$_3$ sample (inset in Figure 5a) pressed from the powder precipitated in an external magnetic field and then dried without mixing. The observed variation in brightness of colored parts the crystallite block was caused by a tilted orientation of its optical axis.

These preliminary results suggest a good potential of this approach to produce relatively big self-supporting samples with oriented LDH crystallites.

Crystallites of the obtained Co$_2$Al–NO$_3$ were observed to be hexagon-shaped (Figure 7), which is typical for the rhombohedral LDHs.

Free precipitation of LDH crystallites from water suspension is known to result in formation of big shapeless particles that consist of hundreds and even thousands of the crystallites [31]. If the LDH particles were dispersed in ethanol, agglomeration was mainly suppressed. Nevertheless, when an external magnetic field was not applied, the LDH crystallites precipitated randomly (Figure 8b,d). The application of pulsed magnetic field promoted a formation of a dense homogenous layer of the flat-laying crystallites on a surface of the substrate (Figure 8a,c). One should notice that owing to a compact arrangement of the LDH crystallites the obtained film was thinner by about a factor of four than that in the case of randomly oriented crystallites.

It has been found that the crystallites that precipitated in an external magnetic field form also some regular structures on the surface of the thin dense LDH film (inset in Figure 8). We associate this phenomenon with orientation effect of a magnetic field on the LDH crystallites—nanoflakes.
during the precipitation process. Our preliminary calculations revealed a highly anisotropic magnetic susceptibility of Co-containing octahedral layers that leads to a predominant space orientation of the LDH nanoflakes in a magnetic field.

![Figure 8](image)

**Figure 8.** SEM images of layers of the Co₂Al–NO₃ LDH crystallites deposited on glass substrates with (a,c) and without (b,d) application of an external magnetic field; cross section views (a,b) and top views (c,d). Inset shows the regular-shape particles of Co₂Al–NO₃ formed on the surface of dense LDH film.

4. Conclusions

The use of high-power sonication for the production of layered double hydroxides emerges as a good option for optimization and improvement of the quality of LDH production.

When high-power sonication was applied to accelerate the processes of the Co₂Al–NO₃ LDH crystallization and the nitrate-to-carbonate anion-exchange, it was possible to reduce the duration of these processes from 4 h to 5 min (crystallization) and from 7 days to 7 min (anion exchange).

The magnetic field assisted approach to the formation of homogeneous oriented layers of deagglomerated crystallites—flakes of Co²⁺-containing LDHs—is promising and deserves further development and optimization. Although crystallites of Co:Al LDHs are typically small and inclined to agglomeration that makes the studies related to their magnetic anisotropy difficult, they can be dispersed and precipitated as ordered layers in an external magnetic field. Semi-transparent samples of 0.15–0.30 mm thick with relatively big blocks (up to 0.5 mm wide) of ordered LDH crystallites are obtained from thick layers precipitated and then pressed into compacts. Thin transparent dense films of Co:Al LDHs are formed from ordered crystallites precipitated from diluted suspensions.

**Author Contributions:** A.N.S. proposed the main idea and wrote the manuscript. D.E.L.V. prepared the samples, performed XRD characterization, TEM and STEM studies. I.M.L and Y.O.S. conducted the optical experiments. E.L.F. and A.V.F. calculated magnetic susceptibility of Co-containing LDH crystallites. Y.G.P. and R.Y.B. justified theoretically the possibility of magnetic field-assisted arrangement of the crystallites. V.V.R. designed the setup for the high-power ultrasound treatment. A.D.S. constructed and calibrated the setup and controlled the sonication-assisted experiments. M.G.S.F. and J.M.V. provided useful discussion and interpretation of the obtained results and coordinated the work.

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