Direct Synthesis of Quaternary Alkylammonium-Capped Perovskite Nanocrystals for Efficient Blue and Green Light-Emitting Diodes

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ABSTRACT: Cesium lead halide nanocrystals (CsPbX₃ NCs) are new inorganic light sources covering the entire visible spectral range and exhibiting near-unity efficiencies. While the last years have seen rapid progress in green and red electroluminescence from CsPbX₃ NCs, the development of blue counterparts remained rather stagnant. Controlling the surface state of CsPbX₃ NCs had proven to be a major factor governing the efficiency of the charge injection and for diminishing the density of traps. Although dodecylammonium halides (DDAX; X = Br, Cl) had been known to improve the luminescence of CsPbX₃ NCs when applied post-synthetically, they had not been used as the sole long-chain ammonium ligand directly in the synthesis of these NCs. Herein we report a facile, direct synthesis of DDAX-stabilized CsPbX₃ NCs. We then demonstrate blue and green light-emitting diodes, characterized by the electroluminescence at 463−515 nm and external quantum efficiencies of 9.80% for green, 4.96% for sky-blue, and 1.03% for deep-blue spectral regions.

In the past five years, colloidal lead halide perovskite nanocrystals (NCs) of APbX₃ composition (A = cesium, formamidinium, or methylammonium; X = Cl, Br, or I) have emerged as a novel class of highly fluorescent semiconductor quantum dots with emission spectra spanning the entire visible range, characterized by narrow emission bands (<100 meV) and near-unity photoluminescence quantum yields (PLQY). They are widely researched toward applications in conventional liquid-crystal displays (LCDs) with red and green fluorescent NCs as downconverters in the backlighting, in the upcoming display technologies such as front-panel LCD architectures, wherein NCs serve as both light filters and fluorescent emitters, or in LED displays exploiting electroluminescence of perovskite NCs. For these display technologies, perovskite NCs offer a compelling suite of attributes: the widest color gamut (due to spectral tunability and narrowband emission) fulfilling the BT.2020 standard exceptionally low production costs, as well as satisfactory restriction of hazardous substances compliance. Other applications of luminescent perovskite NCs may include solid-state lighting, solar cell concentrators, and scintillators for X-ray imaging. These NCs are also increasingly recognized as highly potent single-photon emitters for the design of quantum light sources for quantum information technologies. Emission from single-perovskite NCs is characterized by narrow emission line width, fast radiative decays (10−1000 times faster than in any other colloidal NCs), and long exciton coherence times. In the form of long-range ordered, self-assembled superlattices, these NCs exhibit an accelerated collective mode of emission at cryogenic temperatures, known as superfluorescence, wherein packets of bundled photons are emitted.

Electroluminescence (EL) of layered 2D lead-halide perovskites had been known since the early 1990s. The interest in PL from lead halide perovskites had revived in 2013−2014 (e.g., thin-film MAPbBr₃ LEDs), and these efforts had been
motivated by and had paralleled the use of MAPbX3 films in photovoltaics. Soon thereafter, the LED reports extended to the newcomer colloidal CsPbX3 NCs, as well as their FA and MA NC cousins.

Reduction of structural and electronic dimensionalities and efficient electronic passivation of the surfaces and interfaces—either using colloidal NCs as presynthesized inks for the deposition of the emissive layers or by inducing layering, compositional segregation and gradients, and nanograin formations in situ upon coating from precursors—had been sought as major means of enhancing the EL properties. These structures benefit from the reduction of the exciton diffusion length, increasing exciton binding energy, and oscillator strength. Rapid progress in the development of perovskite-based LEDs emitting in green and red led to the demonstration of the external quantum efficiencies (EQE) exceeding 20%. As a general trend, NC-based perovskite LEDs fall somewhat behind their precursor-deposited thin-film counterparts; for instance, green LEDs based on NCs have thus far yielded lower best EQE value of 16.48% for Cs/FAPbBr3 NCs, while the majority of reports present values well below 10%. Development of blue perovskite LEDs lags far behind those emitting in green and red regions. For the NC-based LEDs the best EQE values are stagnant at ca. 2% for the region of 460–490 nm.

Congreve and co-workers optimized the device architecture for the deep-blue emitting devices by modifying the hole-transporting layer surface with Na-fluorinated ionomer (PFI) and reached 0.5% for LEDs emitting at 465 nm. More recently, the same group introduced Mn2+ doping for improving PLQY of CsPb(Br/Cl)3 NCs, thereby boosting the EQE values to 2.12% at 466 nm. With device brightness of 212 Cd/m2, this is currently the best-performing perovskite LED in the deep blue region.

An alternative approach to blue perovskite LEDs is to form an emissive layer from lower dimensionality (i.e., 2D) perovskite materials with bulky cations such as butylammonium (BA), phenylethylammonium (PEA), or phenylbutylammonium (PBA), which separate few-octahedra-thick lead halide slabs, thereby introducing strong quantum confinement of the excitons and a blue shift of the emission. Jun Xing et al. used similar, in situ formed 2D perovskites such as PEA2A1.5Pb2.5Br8.5 to fabricate LEDs with EQE of 1.5% at 490 nm. Mn2+ doping of such films (PEA2.2Cs0.8MA0.2Pb(1−x)Mn(x)Br3Cl) slightly improved device efficiency to 0.58% in the blue. Such 2D materials generally exhibit good color stability due to the absence of halide segregation, which is a notable issue with mixed-halide perovskites. State-of-the-art results for such materials

Figure 1. (a and b) Representative PL and absorbance (normalized on excitonic peak) spectra of DDAX-stabilized CsPb(Br1−xClx)3 NCs (inset is the photograph of the colloidal solutions of CsPb(Br1−xClx)3 NCs under UV-light excitation, λ=365 nm). (c) X-ray diffraction pattern of CsPbBr3 (green) and CsPb(Br1−xClx)3 (blue) and schematic illustration of the orthorhombic lattice of CsPb(Br1−xClx)3 NCs. The diffractograms are shifted vertically for clarity. (d and e) Transmission electron microscopy images of CsPb(Br1−xClx)3 NCs, for x = 0 and x = 0.24.
show that highly confined (Cs/FA)PbBr₃ NCs formed in situ within the quasi-2D PbABr phase allowed achieving EQE of 9.5% at 483 nm while showing the brightness of 700 cd/m².⁶⁶

The lower performance of green and red NC-based LEDs, compared to the best-performing thin-film counterparts, is commonly attributed to the diverse roles played by long-chain surface capping ligands. On the one hand, the use of long-chain ligands allows for unmatched control over the NC size and shape as well as for ensuring morphological homogeneity within the NC ensemble. These ligands are paramount for the entire life cycle of NCs, from the synthesis of NCs to the retention of the long-term structural and colloidal integrity and for their processing into thin, homogeneous films. On the other hand, controlling and adjusting the inorganic–organic interface of colloidal perovskite NCs had proven to be cumbersome and far more complex as compared to the present state in the surface science of other semiconductor NCs. Structural softness, lattice polarity, and finite solubility in the wide variety of solvents are the factors causing overall lability of perovskite NCs. In this context, highly dynamic and loose binding of the capping ligands, which are thus far most effective in the perovskite NC synthesis [oleic acid (OA) and oleylamine (OLA)], hinders the efficient isolation and purification of NCs and necessitates the presence of the same or alternative ligand in the excess for maintaining the NC stability. This, in turn, imparts the density and other morphological characteristics of the NC films and hence the charge transport through the device. While solution- or solid-state ligand exchange strategies had been highly successful in mitigating these challenges with conventional semiconductor NCs (typically Cd-, Pb-, and other metal chalcogenides), little progress had been achieved with perovskite NCs. In the context of LED research, the most effective approach has been to use didodecyl(dimethylammonium bromide (DDAB) for a postsynthetic surface treatment⁶⁷–⁷² or as an additive during the synthesis.⁴⁵,⁷³ Comonstituent passivations of undercoordinated Pb-atoms with bromide anions and strong affinity of the DDA-cation to the surface A-site give rise to both electronic passivation and hence near-unity PLQY values, as well as improved stability and processability, as had been thoroughly examined experimentally and computationally by us and others.⁵–⁷⁷ Pan et al. applied DDAB-treatment to CsPb(Br/Cl)₃ NCs in solutions, reaching 1.9% EQE at 490 nm; however, rather low brightness of 35 Cd/m² was reported.⁶⁷ Combined application of both DDAB and its chloride analog (didodecyl(dimethylammonium chloride, DDAC) had been shown to yield EQE values of 0.86% at 480 nm and 0.1% at 458 nm.⁷²

In striving to harness the beneficial effects of DDAX in the most direct manner, we present a novel synthesis of uniform, cuboid CsPb(Br₁₋ₓClₓ)₃ NCs (x = 0–0.39), wherein DDAB and/or DDAC are used as a sole ligand system, entirely eliminating the use of OA and OLA. The PL peak position of CsPb(Br₁₋ₓClₓ)₃ NCs can be accurately adjusted in the range of 462–510 nm, whereas PLQY reaches 80%. NCs can be well-purified and consequently allow for fabrication of uniform and compact, thin films. Through judicious optimization of the LED device architecture, high EQE values of 9.8%, 4.96%, 3.5%, 2.25%, and 1.03% have been obtained for EL peak positions at 512, 502, 490, 476, and 463 nm, respectively. Highly luminescent CsPb(Br₁₋ₓClₓ)₃ NCs with adjustable chloride content (represented by x as a molar proportion of chloride) were synthesized using hot-injection methodology with DDAX as a sole long-chain (C₁₂) capping ligand. The synthesis, outlined in Figure 1, was carried out in mesitylene as a solvent and under nitrogen atmosphere. PbBr₃ was first solubilized in mesitylene by mixing with DDAB or (DDAB–DDAC) and quickly heated to 150–160 °C to obtain a transparent colorless solution. PbBr₃ alone is not soluble in mesitylene; however, it forms lead-bromide anionic species (presumably PbBr₃⁻ and the like) when combined with DDAB at elevated temperatures (>100 °C). Recently, Song et al. reported the use of the quaternary ammonium salts such as tetaoctylammonium bromide (TOAB) in the synthesis of CsPbBr₃ NCs at room temperature, followed by introducing DDAB ligands on the surface of the formed NCs.⁴⁸ While TOAB aids in solubilizing PbBr₃ at room temperature, it is too bulky for binding to the NC surface. We find that for the synthesis at elevated temperatures DDAB alone can play both roles (solubilizer and surfactant). As a source of Cl ions, DDAC was found to be more convenient than the addition of PbCl₂; the latter exhibits very low solubility in nonpolar solvents. A separate Cs⁺ stock solution was prepared by dissolving Cs₂CO₃ in disooctylphosphinic acid and toluene. The Cs solution was then swiftly injected into the first solution. An instant nucleation of NCs is seen as a rapid yellow-green coloration. In this synthesis, we completely omitted OA and OLA as capping ligands, which dynamically and loosely bind to the surface of lead halide perovskite NCs and easily desorb.⁷⁸ Earlier studies indicated that oleate ligands are mostly absent on the surface of such NCs,⁷⁴,⁷⁵,⁷⁹ and the main role of OA is the protonation of OLA. The trouble with OLA cation is that it rapidly loses its proton and desorbs from the NC surface. In contrast, DDA is a quaternary cation with a static positive charge, and hence, no acid is needed in the reaction mixture for adjusting the pH value. DDAX-stabilized NCs sustain two washing steps by the repetitive destabilization with the moderately polar solvent (acetone–toluene mixture) and redispersion in cyclohexane, as can be judged from the retention of optical characteristics (Figure S1). We note that DDAX-coated NCs are not dispersible in toluene, which is a typical solvent of choice for NCs capped with longer-chain OLA and OA ligands. For the subsequent experimentation with LED devices, we have chosen to focus on weakly confined 9–10 nm large NCs and adjust their band gap energy by the halide composition. A detailed study on the size- and shape-control in this DDAX-based synthesis, as well as on the specifics of surface chemistry using NMR spectroscopy and molecular dynamics calculations, is underway and will be published elsewhere.

Elemental analysis of the thin films of purified NCs evidences that the mixing ratios of DDAB:DDAC directly translate into the composition of CsPb(Br₁₋ₓClₓ)₃ NCs (Table S1). The fact that the halide composition of the NCs reflects an overall halide ratio in the reactions mixture stands in good agreement with OLA/OA-based preparation of CsPb(Br₁₋ₓClₓ)₃ NCs (composition adjusted by PbCl₂/PbBr₂)¹⁶,⁸⁰ and other synthesis methodologies,⁵² including postsynthetic compositional adjustments by the anion-exchange reactions.⁸¹,⁸² Notably and in contrast, FAPb(Br₁₋ₓClₓ)₃ NCs are not stable and separate into two monohalide phases,⁸³,⁸⁴ as demonstrated by both the conventional synthesis methodologies and thorough screening of reaction parameters by the microfluidic-based high-throughput experimentation,⁸⁵ hence, they are not suited for engineering blue LEDs.
The composition-tunable PL and absorption spectra of purified CsPb(Br1−xClx)3 NCs (x ≤ 0.39, Figure 1 and Table 1) span the violet to green spectral region (462–510 nm), with blue shift governed by the quantity of incorporated Cl. Owing to the weak confinement regime at perovskite NC sizes of 8–10 nm35,86 and hence minimal impact of the size-distribution on PL full width at half-maximum (fwhm), a rather narrowband emission had been attained (fwhm = 18–20 nm).

Purified NCs show PLQYs of 80, 62, 48, 32, 25% for PL emission bands centered at 510, 499, 487, 476, and 462 nm, respectively (Figure S2). Colloidal solutions of CsPb(Br1−xClx)3 NCs obtained by the DDAX approach are stable for months without losing their PLQYs and colloidal stability.

The XRD pattern in Figure 1c confirms the orthorhombic perovskite structure of CsPb(Br1−xClx)3 NCs with no noticeable presence of other phases. All main diffraction peaks in the case of Br/Cl-containing NCs were shifted toward larger angles compared to the ICSD-#97851-pattern (orthorhombic phase of CsPbBr3), because of the lattice contraction induced by the smaller radius of the Cl ions, thereby also confirming the formation of the homogeneous solid solution. The TEM image reveals that CsPbBr3 and CsPb(Br1−xClx)3 NCs exhibit uniform cuboid morphology with a crystal size of ~9 nm (Figure 1d,e). The colloids and corresponding drop-cast films from CsPb(Br1−xClx)3 NCs were also characterized by the TRPL. All samples showed biexponential decay lifetimes in solution with intensity averaged lifetime values of 10.5, 4.6, 10.4, 7.7, and 20.7 ns (for x-values from 0.39 to 0), from bluest to greenest samples respectively (Figure S3). Drop-cast films generally exhibited shorter lifetimes of 8.5, 1.95, 2.8, 2.84, and 2.34 ns for the same samples, correspondingly.

Initial electroluminescence tests had been conducted on Cl-free samples (CsPbBr3 NCs) in order to benchmark the utility of the new synthesis and compare to the extensive prior art on CsPbBr3 NC LEDs. The best LED characteristics were attained when using 15 nm layers of B3PYMPM (4,6-bis(3,5-di(pyridin-3-yl)phenyl)-2-methylpyrimidine) or 3TPYMB (3,3′,3″-[borylidynetris(2,4,6-trimethyl-3,1-phenylene)]tris[pyridine]) as hole-blocking layers (HBLs), in

### Table 1. Summary of the Material and Device Performance Parameters of LEDs with CsPb(Br1−xClx)3 NCs of Various Halide Compositions

| Sample, CsPb(Br1−xClx)3 | DDAC/DDAB ratio in the synthesis | PL peak(a) (nm) | PLQYx | best EQE | EQE@100 Cd | best luminance (Cd/m2) | EL peak position (nm) |
|--------------------------|---------------------------------|-----------------|--------|----------|-------------|------------------------|----------------------|
| x = 0.39                 | 1:0                             | 462             | 25%    | 1.03%    | 0.23%       | 193                    | 463                  |
| x = 0.24                 | 6.5:3.5                         | 476             | 32%    | 2.25%    | 1.07%       | 678                    | 476                  |
| x = 0.19                 | 6.1:3.9                         | 487             | 48%    | 3.5%     | 2.1%        | 2,063                  | 490                  |
| x = 0.11                 | 3.7:6.3                         | 499             | 62%    | 4.96%    | 3.94%       | 7,600                  | 502                  |
| x = 0                        | 0:1                             | 510             | 80%    | 9.8%     | 7.3%        | 34,700                 | 512                  |

x In solution.

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Figure 2. Performance of green-emissive LED with DDAB-capped CsPbBr3 NCs. (a) Energy levels of active layers of the device vs vacuum. (b) Current density and luminance vs applied bias. (c) EL spectra for various driving voltages. (d) External quantum efficiency vs current density through the active area.

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ACS Energy Letters

DOI: 10.1021/acsenergylett.9b01915

ACS Energy Lett. 2019, 4, 2703–2711
combination with a 45 nm layer of TPBi (1,3,5-tris(1-phenyl-1H-benzimidazol-2-yl)benzene) as an electron transporting layer (ETL) and pTPD (poly[N,N′-bis(4-butylphenyl)-N,N′-bisphenylbenzidine]) as a hole-transporting layer (HTL). Figure S4 evidences the beneficial effects of HBLs by comparison with HBL-free devices. The best-efficiency devices were those employing a 15 nm layer of B3PYMPM as an HBL and a 45 nm layer of TPBi as an ETL. These devices presumably benefit from hole localization in the emissive layer (at TPBi), thereby balancing the barrier existing for the injection of holes. This device architecture, together with the judicious optimization of the NC purification (the choice of solvents and nonsolvents and their amounts, as well as number of purification steps etc.), yielded a champion green device, with characteristics shown in Figure 2. Specifically, the EL peak centered at 515 nm was characterized by EQE values of up to 9.8%. High EQE values are satisfactorily retained at higher current densities, allowing for obtaining brightness of 34 700 \text{Cd/m}^2. Other reports on LEDs employing CsPbBr\textsubscript{3} NCs with DDAB treatment have presented EQE values of 7−9.7% and brightness of 2269−15000 \text{Cd/m}^2.\textsuperscript{70,87}

Blue-emitting LEDs were fabricated with the device structure proposed by Gangishetty et al., where a TFB (poly[(9,9-dioctylfluorenyl-2,7-diyl)-co-(4,4′-(N-(4-tri-butylphenyl) diphenylamine)]) layer covered with Naion perfluorinated ionomer (PFI) is used as an HTL for improved hole injection.\textsuperscript{60} Figure S5 shows device performance of our LEDs based on DDAX-stabilized CsPb(Br\textsubscript{1−x}Cl\textsubscript{x})\textsubscript{3} NCs with EL peak at 480 nm. While EQE of \textasciitilde0.57% is a modest improvement in comparison to ref 60 (0.44% at 481 nm, OLA/OA-synthesized NCs), the brightness was approximately 10-fold higher and reached 2500 \text{Cd/m}^2 (vs 212 \text{Cd/m}^2 in ref 60). We should also note that we were not able to fully suppress the EL from the TFB layer (at \textasciitilde430 nm) by the application of PFI, presumably because of the suboptimal thickness of the perovskite NC layer and hence electron injection into TFB.

Figure 3. (a and b) SEM cross-sectional image and corresponding schematic of the blue LED stack. (c) Work function of electrodes and band gap values of the different materials used in the LED stack, energies vs vacuum. (d) Current density and luminance of the devices with various EL peak positions as a function of voltage. (e and f) EQE of the devices with various EL peak positions as a function of current density and luminance.

Figure 4. (a) EL spectra of the LEDs. Legend reveals corresponding EL peak position. Inset shows photo of the EL of the operating device with Empa logo. (b) Commission Internationale de l’Eclairage CIE 1931 color diagram. Black triangle represents BT.2020 (Rec2020) color coordinates. Red dots mark coordinates of LEDs presented in this work.
Additional device structure optimization was made by using a two-step hole injection with both pTPD and another HTL material with a deeper highest occupied molecular orbital level, namely, PVK (poly-[9-vinylcarbazole]). The best device structure for the blue LEDs was the following (Figure 3a–c): ITO/PEDOT:PSS/pTPD/PVK/NCs/B2PyMPM (15 nm)/TPBi (45 nm)/LiF (1.2 nm)/Al (100 nm). Current-density curves reveal low leakage currents of the presented device structure (Figure 3d). The turn-on voltages are in the range of 3–3.4 V, which is still higher than the emissive material band gaps (2.47 – 2.68 eV). EQE versus current-density and luminance curves of the devices presented in Figure 3e,f show efficiency drops at higher currents/luminance. The best EQE values are 4.96%, 3.5%, 2.25%, and 1.03% for the devices with a EL peak at 502, 490, 476, and 463 nm, respectively (Figure S6). Lower efficiency in a deep-blue spectral region might be associated with the lower PLQYs of Cl-containing NC perovskites.

All devices showed high color purity and narrow EL spectra without any visible emission from the transport layers (Figure 4a), indicating that the layer of NCs is the preferable recombination zone in the devices. Chromaticity coordinates achieved in the devices presented here are presented on Figure 4b and Table S2, with one of the devices reaching the recommended blue color for BT.2020.52 Table 1 summarizes the synthesis conditions and optical and optoelectronic properties of CsPb(Br1_xCl_x)3 NCs and respective LEDs.

We further tested the device stability and evolution of the electroluminescence. Increased driving voltage does not cause changes in the EL peak position (Figure S7). The constant-current driving under initial brightness of 100 Cd/m2 of the device with CsPb(Br0.76Cl0.24)3 NCs results in the EL intensity drop to 20% of the initial value within 26 s, which, although fast, is comparable to the best blue-emitting perovskite LEDs made from both the NCs and quasi-2D perovskites.61,64,66 We also observe red shift of the electroluminescence spectrum (Figure S8). Such color instability is commonplace in perovskite devices and is caused by halide ion migration during the device operation.88–91 In Br/I systems, the halide segregation is readily observed under illumination and without applied electric bias and is commonly assumed to occur under the action of the local electric field induced by free and trapped charge carriers.2,93 In the case of electrical charge injection, this effect might be more pronounced because of possible charge imbalance and higher electrical field. While reported EQEs of perovskite LEDs are rapidly increasing, the stability of devices remains a major obstacle. Further studies are required to unveil the factors governing the long-term stability, which may include degradation of the charge-transporting layers or the perovskite layer. We tend to assume that the fundamental issue lies with the facile electrochemical reduction of Pb2+ to metallic Pb. If true, this degradation path should be more pronounced for higher electronegativities of halide, e.g. for chlorides.

In summary, we reported a direct synthesis of long-chain quaternary ammonium-capped CsPb(Br1-xClx)3 NCs, namely employing DDAX as the sole ligand in the hot-injection synthesis. With its C12-hydrocarbon chains, this ligand is considerably shorter than OLA and OA (C18), favoring the application of such NCs in LEDs. Green-emissive LEDs exhibit a peak EQE of 9.8% and brightness of 34700 Cd/m2 and are, to the best of our knowledge, the most efficient devices based on CsPbBr3 NCs. Application of double hole-transporting layers and additional hole-blocking layers afforded efficient blue LEDs (EQE = 1–5% at EL peaks of 460–500 nm). Future work should focus on the maximization of the PLQY values of the NCs before integration into the LEDs. Another highly promising venue is solid-state chemical treatment, e.g. by dipping the films of DDAX-capped NCs into solutions of diverse chemical species, which could allow for passivation of the trap states.

**ASSOCIATED CONTENT**

* Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acsenergylett.9b01915.

Experimental methods; materials; green LED device structure optimization; spectra of the blue-emissive LEDs with various driving voltages; device stability test; color coordinates of the LEDs (PDF)

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**Notes**

The authors declare no competing financial interest.

**ACKNOWLEDGMENTS**

Authors are thankful to Dr. Mario Ochoa Gomes and Dr. Yaroslav Romanyuk for access to the TRPL measurement setup, Dr. Oleksii Liashenko for help with SEM imaging, and Dr. Frank Krumeich for SEM-EDS compositional analysis. The authors are grateful to the Scientific Center for Optical and Electron Microscopy (ScopeM, ETHZ) and Empa Electron Microscopy Center for use of their instruments. This project has received funding from the European Research Council (ERC) under the European Union’s Horizon 2020 research and innovation program (Grant Agreement No. 819740), project SCALE-HALO, from Swiss Federal Laboratories for Materials Science and Technology (Empa), and Swiss Federal Institute of Technology in Zurich (ETH Zurich).

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