

BOston Neonatal Brain Injury Dataset for Hypoxic Ischemic Encephalopathy (BONBID-HIE): Part I. MRI and Manual Lesion Annotation

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ABSTRACT

Hypoxic ischemic encephalopathy (HIE) is a brain injury that occurs in 1 ~ 5/1000 term neonates. Accurate identification and segmentation of HIE-related lesions in neonatal brain magnetic resonance images (MRIs) is the first step toward predicting prognosis, identifying high-risk patients, and evaluating treatment effects. It will lead to a more accurate estimation of prognosis, a better understanding of neurological symptoms, and a timely prediction of response to therapy. We release the first public dataset containing neonatal brain diffusion MRI and expert annotation of lesions from 133 patients diagnosed with HIE. HIE-related lesions in brain MRI are often diffuse (i.e., multi-focal), and small (over half the patients in our data having lesions occupying <1% of brain volume). Segmentation for HIE MRI data is remarkably different from, and arguably more challenging than, other segmentation tasks such as brain tumors with focal and relatively large lesions. We hope that this dataset can help fuel the development of MRI lesion segmentation methods for HIE and small diffuse lesions in general.

Background & Summary (700 words maximum)

Accurate identification of brain lesion injuries in neonatal brain magnetic resonance images (MRI) [1, 2, 3] is crucial to improve clinical care of neonates with hypoxic ischemic encephalopathy (HIE), a brain disease that occurs in around 1 ~ 5/1000 term-born infants at birth [4, 5]. HIE affects around 200,000 term-born neonates every year worldwide [4, 5], costing about \$2 billion/year in the US alone, let alone family burdens. Therapeutic hypothermia, the current clinical treatment of HIE, can reduce mortality and morbidity. Nevertheless, around 60% of patients still die or develop neurocognitive deficits by 2 years of age. MRI is used in over 50% of all the >100 ongoing HIE-related clinical trials worldwide [6], for evaluating treatment effects [7, 8, 9], and helping discover clinical [10, 11, 12], biochemical [10, 13, 14, 15], and serum [16, 17, 18] biomarkers. Accurate identification of brain lesions in neonatal brain MRIs [1, 2, 3] is needed for disease prognosis, a better understanding of the neural basis of disease progression, and more timely evaluations of therapeutic effects.

HIE lesions are often diffuse (i.e., multi-focal), and small; hence, algorithms that have shown great promise in segmenting big and focal lesions, such as brain tumors and acute strokes, often encounter challenges when directly applied to MRIs of HIE patients. Indeed, many (over half) patients had lesions occupying <1% of brain volume, as shown in Figure 1. As a result, the segmentation accuracy measured by the Dice overlap with U-Net [19] and other state-of-the-art machine/deep learning algorithms on HIE remains at around 0.5 [20], whereas Dice is over 0.8 when segmenting brain tumors [21, 22].

A major hurdle in developing algorithms for small diffuse lesions, such as HIE lesions, is the lack of public data. Public data with expert annotations of lesions have fueled the advancement of machine learning algorithms to segment brain tumors [23], stroke lesions [24], multiple sclerosis lesions [25, 26], and numerous other diseases in the brain or other organs [27, 28]. However, to date, there is no public MRI data with expert annotations available for HIE lesions.

We present BOston Neonatal Brain Injury Dataset for Hypoxic Ischemic Encephalopathy (BONBID-HIE), an open-source, comprehensive, and representative MRI dataset for HIE. This paper introduces the first part of the BONBID-HIE data. This release contains raw and derived diffusion parameter maps, as well as manually-annotated lesion masks, for 133 HIE patients.

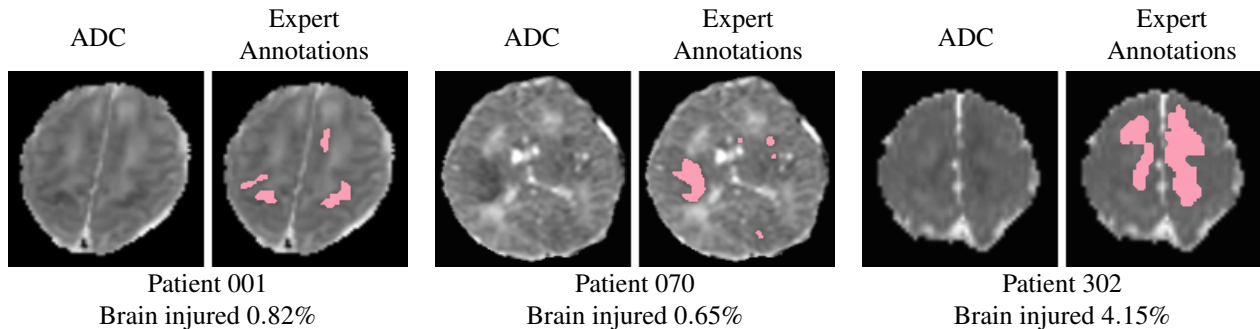


Figure 1. Lesions associated to hypoxic-ischemic encephalopathy (HIE) are typically diffuse (i.e., multi-focal) and small. Here we show two representative images for 3 HIE patients. For each patient, in the left panel: apparent diffusion coefficient (ADC) maps that are clinically used to identify HIE lesions; in the right panel: manually-annotated lesions (shown in pink) overlaid on the ADC map. We listed the percentage of the whole brain volume being injured at the bottom (i.e., lesion volume divided by the whole brain volume).

38 Our data was from Massachusetts General Hospital. It includes MRIs from different scanners (Siemens 3T and GE 1.5T),
 39 different MRI protocols, and from patients of different races/ethnicities and ages (0-14 days postnatal age). Part I of our
 40 data release (this paper) focuses on lesion detection, while Part II (a follow-up paper) will focus on clinical, treatment, and
 41 neurologic outcome data for further developing prognostic biomarkers.

42 Methods

43 This work was approved by the Institutional Review Boards (IRBs) at Massachusetts General Hospital (MGH) and Boston
 44 Children's Hospital (BCH). Figure 2 illustrates the overall data archiving process.

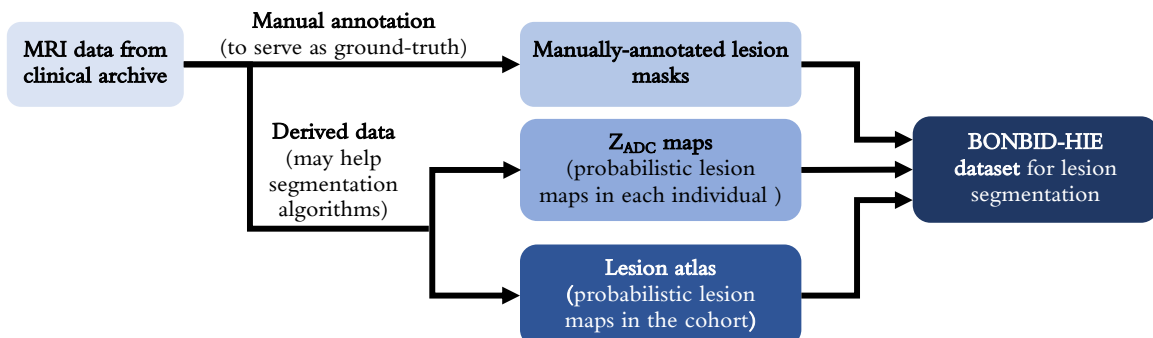


Figure 2. A schematic diagram showing the steps performed on the BONBID-HIE data for release.

45 Retrospective Data Collection

46 Data was retrospectively collected from MGH. Inclusion criteria were: (1) term-born (at physician discretion) (2) clinical
 47 diagnosis of HIE; (3) initially treated at MGH between 2001 and 2018; (4) no comorbidities such as hydrocephalus or congenital
 48 syndromes; and (5) high-quality MRI acquired in Day 0-14 after birth (visually checked by RW, AF, YO). Exclusion criteria
 49 were: (1) excessive motion artifacts or missing images; (2) secondary HIE diagnosis to a primary perinatal stroke.

50 Clinical characteristics and demographic information were retrospectively gathered from the electronic health records
 51 (EHRs). The clinical variables included maternal information during pregnancy and delivery, as well as infant information.
 52 More detail can be found in the "Data Records" Section.

53 MRI data was downloaded from MGH Radiology Department clinical archives using the mi2b2 search engine [29]. MRIs
 54 were acquired on either a GE 1.5T Signa scanner (N=52, scanned during 2001-2012), or, a Siemens 3T TrioTim or PrismaFit
 55 scanner (N=81, scanned during 2012-2018). Diffusion tensor sequences on all scanners had the protocol as follows: Time
 56 of Repetition (TR)=7500 – 9500ms, Time of Echo (TE)=80 – 115ms, and b=1000 s/mm². The GE scanner had resolution
 57 1.5 × 1.5 × (2.0 – 4.0)mm³ and (6 - 60) diffusion directions, while the Siemens scanner had a resolution 2 × 2 × 2mm³ and (25

58 - 60) diffusion directions. Apparent diffusion coefficient (ADC) maps were directly generated by the scanners (with Syngo
 59 software for Siemens scanners [30], and with the Advantage Windows Workstation for GE scanners [31, 32]).

60 MRI Pre-processing

61 Besides the raw NIfTI image as converted from the DICOM files, we also generated several processed images. The pre-
 62 processing steps included: N4 bias correction [33], field of view normalization [34], multi-atlas skull stripping for the ADC
 63 maps [35], and deformable registration of each patient's ADC map to a normative 0-14 day neonatal brain ADC atlas [36], by
 64 the Deformable Registration via Attribute Matching and Mutual-Saliency weighting (DRAMMS) software [37], which has
 65 been extended-validated for lifespan ages in various MRI sequences [38]. This normative ADC atlas was constructed from
 66 ADC maps of 13 healthy individuals acquired 0-14 days after birth (Figure 3a) with our extensively-validated MRI analysis
 67 pipeline [39, 38, 35, 34]. All software packages used in this pre-processing pipeline are publicly available and have been
 68 validated in processing both research and clinical MRI scans across ages [40, 41, 42, 43].

69 Expert Annotation of Lesions

70 HIE lesions were manually annotated as a binary mask on the 3D ADC maps in the patient's raw image space, using the
 71 MRICroN software. ADC maps are clinically used as the standard images to identify HIE abnormalities [44, 45]. The
 72 annotations were done by a trained physician (YS; >3 years of experience) according to the neuroradiology reports that were
 73 generated as part of the clinical flow. The annotations started from the axial slice and were subsequently modified in the
 74 coronal and sagittal planes for the 3D integrity of lesion regions. Uncertainties occurred in 27 patients and were resolved by the
 75 consensus of three pediatric neuroradiologists (CJ, SS, and PEG; >5, >5, and >20 years of experience).

76 Generation of Z_{ADC} Maps for Each Patient

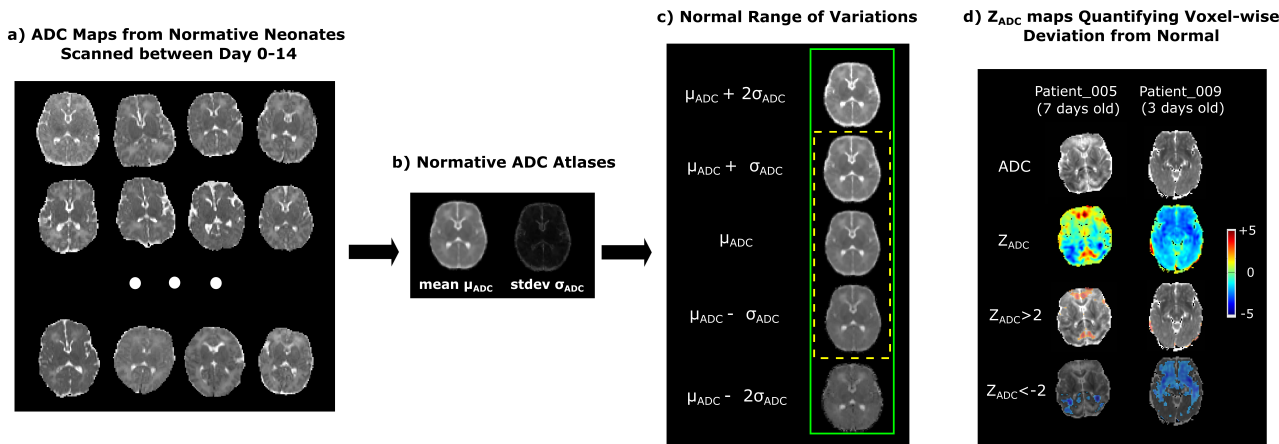


Figure 3. The generation and concept of Z_{ADC} maps. (a) Examples of ADC maps from normative subjects, which were warped into the same space using unbiased group-wise DRAMMS registration to generate (b) the mean and standard deviation ADC atlases. (c) One or two standard deviations above and below the mean ADC atlas define the normal ranges of voxel-wise ADC variations. (d) Our novel Z_{ADC} map quantifies voxel-wise deviations from the mean ADC map in (b). The cool/warm colors in (d) represent voxels with ADC values lower/higher than the mean ADC at the same anatomic location, according to the scale bar on the right.

77 Neuroradiologists identify acute brain injury from HIE as regions with low ADC values. Low ADC values represent a
 78 reduced water diffusion or blood supply [6]. However, a dilemma is, what ADC value is considered abnormally low versus
 79 just low within the normal variation? The normal variations of ADC values differ across brain regions [36, 46], making this
 80 question difficult even for experienced neuroradiologists. For example, a voxel with an ADC value of $800 (\times 10^{-6} mm^2/s)$ may
 81 be considered normal at one brain region, whereas another voxel with an ADC value of $900 (\times 10^{-6} mm^2/s)$ may be considered
 82 lesioned at another brain region, if the normal ranges of ADC variations in the two brain regions are 700-900 and 950-1100
 83 (same unit), respectively.

84 To address this dilemma, we have developed Z_{ADC} maps to normalize and make ADC values comparable across brain voxel
 85 locations [40]. First, a normative ADC atlas was generated from scans of healthy neonates (Figure 3a). This atlas quantifies the
 86 mean ADC values and standard deviation at every voxel [47] (Figure 3b), and hence the normal range of variations at each

voxel (Figure 3c). Then, we converted each patient's ADC map (first row, Figure 3d) into a Z_{ADC} map (second row, Figure 3d). The Z_{ADC} maps compared the patient's ADC value at each voxel to the normal variations at the corresponding voxel in the atlas. In a nutshell, Z_{ADC} maps quantify how many standard deviations away a patient's ADC value at a voxel is from the normal mean at the same anatomical location.

Specifically, a deformation (D) was computed, which mapped every voxel x in the patient's ADC map to its anatomically-corresponding location $D(x)$ in the atlas space. The normal range of ADC variation per voxel was defined by the mean and standard deviation denoted for that voxel across all healthy neonates. Finally, the patient's ADC value I_x at voxel x was converted to a Z value: $Z_x = (I_x - \mu D(x)) / \sigma D(x)$. We calculated the Z_{ADC} map, which resides in the patient's raw ADC image space, for each patient. This offers an option for developing anatomy-aware lesion segmentation algorithms [48].

96 Construction of Statistical Lesion Atlases for the Cohort

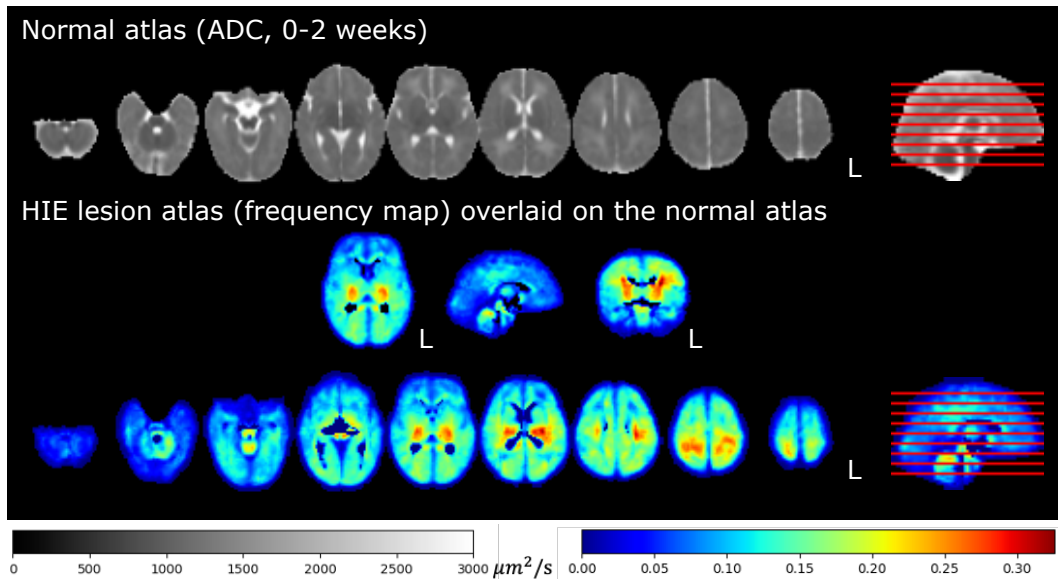


Figure 4. Statistical lesion atlas quantifying the voxel-wise lesion frequency in our cohort of N=133 patients in the normal 0-14 days ADC atlas space.

The same deformation field that was computed by the non-rigid registration from the patient's skull-stripped ADC map to the normal ADC atlas was used to transform the binary brain lesion maps of each patient into the normal neonatal ADC atlas space [37]. The transformed binary lesion masks were then summed and divided by the total number of patients at each voxel. This led to a statistical lesion atlas that quantifies voxel-wise frequency, or probability, of HIE lesions in our cohort, as illustrated in Figure 4.

102 Data Records

103 Dataset Characteristics

Table 1A lists the demographics and clinical characteristics of mothers and neonates. Maternal information includes demographics (age at delivery, race), birth mode (C-section or vaginal), and complications during pregnancy and delivery. Neonatal information includes demographics (age at MRI scan, gestational age at birth, birth weight, head circumference, sex), birth conditions (1/5/10-minute APGAR scores, lowest pH value in umbilical cord), treatment (hypothermia or not), and complications in the neonatal intensive care unit (NICU), including seizure (yes/no), length of stay (in days), the use of endotracheal tube (ETT, yes/no), and the administration of total parenteral nutrition (TPN, yes/no). In each row, we also listed the number of patients who had such information available.

Table 1B quantifies the distribution of the absolute lesion volumes (in mm^3) and relative lesion volume (percentage of the brain being injured). Here, the relative lesion volume was calculated by the volume of the expert-annotated lesion regions divided by the algorithm-extracted whole brain volumes [34, 35]. The median lesion volume accounted for 0.63% of the whole brain volume. This confirms that over half of the patients had less than 1% of the brain being injured. Table 1C further calculates the distribution of the relative lesion volumes (by the percentage of the brain being lesioned). The absolute and

relative volumes were both computed in the patient's raw ADC image space. The minimum lesion was 0 mm^3 , which is a common issue in HIE – mild HIE cases may not show explicit lesions in neonatal MRIs [49, 50, 51].

Figure 5 shows the ADC map, Z_{ADC} map, and expert annotations of example patients with different HIE lesion percentages. Two patients are shown in each of the four groups: those with lesions occupying <1% (upper left panel), 1-5% (upper right panel), 5-50% (lower left panel), and 50-100% (lower right panel) of the whole brain volume. Overall, around 1 in 2 (55.64%) patients had HIE lesions occupying less than 1% of their brain volume, and 3 in 4 patients (75.19%) patients had lesions occupying less than 5% of their brain. This confirms that HIE lesions detectable in the diffusion MRI in our cohort are often small.

124 Data structure and file formats

All medical imaging files were exported from the Picture Archiving and Communication System (PACS) and converted into the NIfTI format. Segmentation masks created by expert annotations were also saved in NIfTI format. Corresponding scanner metadata from the Digital Imaging and Communications in Medicine (DICOM) header in the .json file format is provided with the datasets. All data in the BONBID-HIE dataset was separated into a training dataset ($n=89$) and a test dataset ($n=44$). Both the training and test sets contain data from both scanners (GE 1.5T Signa and Siemens 3T Trio). The split between the training and test data set has been performed (RB, YO) so that both sets include a similar variance of HIE lesion patterns as shown in Table 1C.

Table 1. Cohort characteristics (N=133)

A. Demographics and Clinical Characteristics		
Maternal Information		
Maternal age at delivery (years)	29.5 ± 6.7	N=133
Race	White (43), Black or African American (7), Hispanic or Latino (15), Multi Race (5), Unknown (57), Other (6)	N=133
Delivery	C-section (78), Vaginal (55)	N=133
Antepartum hemorrhage	Yes (29), No (104)	N=133
Thyroid dysfunction	Yes (5), No (128)	N=133
Pre-eclampsia	Yes (9), No (124)	N=133
Fetal decels	Yes (72), No (61)	N=133
Shoulder dystocia	Yes (8), No (125)	N=133
Chorioamnionitis	Yes (20), No (108)	N=133
Emergency c-section	Yes (69), No (58)	N=133
Neonatal Information		
Age at scan (days)	3.9 ± 2.7	N=133
Gestational age at birth (weeks)	39.1 ± 1.9	N=133
Birth weight (g)	3321.9 ± 615.9	N=133
Infant head circumference (cm)	34.2 ± 1.4	N=85
Sex	Male (74), Female (59)	N=133
1-minute APGAR scores	1.9 ± 1.7	N=133
5-minute APGAR scores	4.2 ± 2.3	N=132
10-minute APGAR scores	5.3 ± 2.1	N=118
Lowest pH value in umbilical cord	7.00 ± 0.20	N=129
Therapeutic hypothermia before MRI?	Yes (86), No (47)	N=133
Endotracheal tube (ETT) in NICU	Yes (78), No (47)	N=125
Total parenteral nutrition (TPN) in NICU	Yes (111), No (21)	N=133
Seizures NICU	Yes (64), No (69)	N=133
Length of stay in NICU (days)	12.10 ± 9.92	N=133
B. Lesion Characteristics (N=133)		
Whole-brain lesion volume – minimum	0 mm^3 (0% of the brain injured)	
Whole-brain lesion volume – 25th percentile	441.96 mm^3 (0.10% of the brain injured)	
Whole-brain lesion volume – median	2765.63 mm^3 (0.63% of the brain injured)	
Whole-brain lesion volume – 75th percentile	24264.27 mm^3 (4.86% of the brain injured)	
Whole-brain lesion volume – maximum	412120.00 mm^3 (82.59% of the brain injured)	
C. Number of Patients by Percentage of Lesions in the Brain (N=133)		
[0%, 1%) of the brain being injured	55.64% (N=74)	
[1, 5%) of the brain was injured	19.55% (N=26)	
[5, 10%) of the brain was injured	5.26% (N=7)	
[10, 20%) of the brain was injured	4.51% (N=6)	
[20, 50%) of the brain was injured	8.27% (N=11)	
[50, 100)% of the brain was injured	6.77% (N=9)	
D. Scanner (N=133)		
GE 1.5T	39% (N=52)	
Siemens 3T	61% (N=81)	

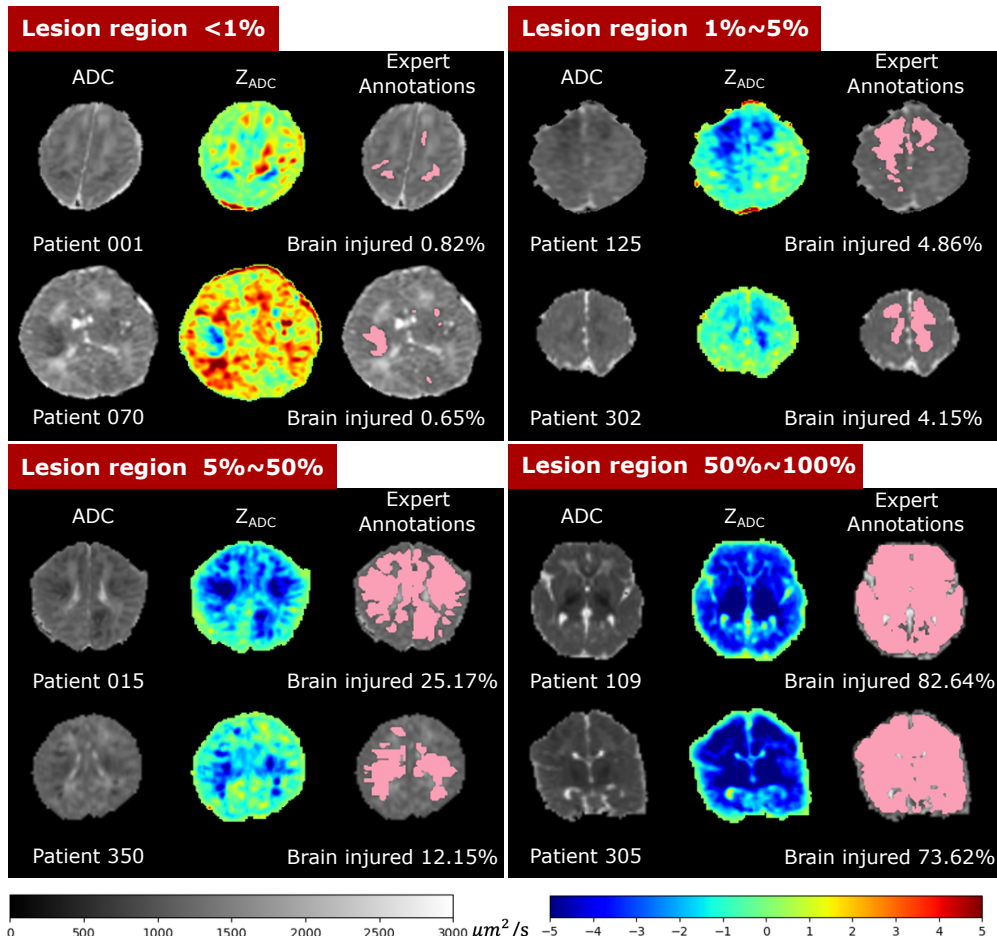


Figure 5. Visualization of patients with different lesion percentages. In every patient, the left image is the ADC map (skull stripped) with range of ADC values designated by the gray scale bar, the middle is the computed Z_{ADC} map with range of Z-scores designated by the rainbow scale bar, and the right image is the expert-annotated lesion regions (pink) overlaid on the ADC map. Percentages of injury were calculated by the volume of the expert-annotated lesion regions divided by algorithm-extracted whole brain volumes.

132 The data is organized in the format shown in Figure 6. BONBID-HIE provides, per patient: (i) 0ADC: raw defaced ADC
 133 maps; (ii) 1ADC_ss: skull stripped ADC map; (iii) 2Z_ADC: Z_{ADC} map; (iv) 3LABEL: expert lesion annotations; and (v)
 134 clinical data: clinical variables as written in Table 1A. There is also (vi) Atlases: a folder for the normal and lesion atlases; (vii)
 135 a readme.txt file: a text file to provide information on this data organization; and (viii) the license file of the BONBID-HIE
 136 dataset.

137 Data repository and storage

138 All training data has been made publicly available under the CC BY NC ND license (<https://creativecommons.org/licenses/by-nc-nd/2.0/>), allowing academic use with credit, prohibiting commercial use without owner's
 139 permission, and disallowing derivation or adaption of data). This dataset is also used for the HIE lesion segmentation challenge
 140 in Medical Image Computing and Computer Assisted Intervention (MICCAI) 2023 annual conference in October 2023.
 141 Further information about the HIE challenges, including data storage and download, can be found under our homepage
 142 (<https://hiechallenge.github.io/>). As such, the expert lesion annotation of lesions in the testing sub-cohort will
 143 be released after the challenge.

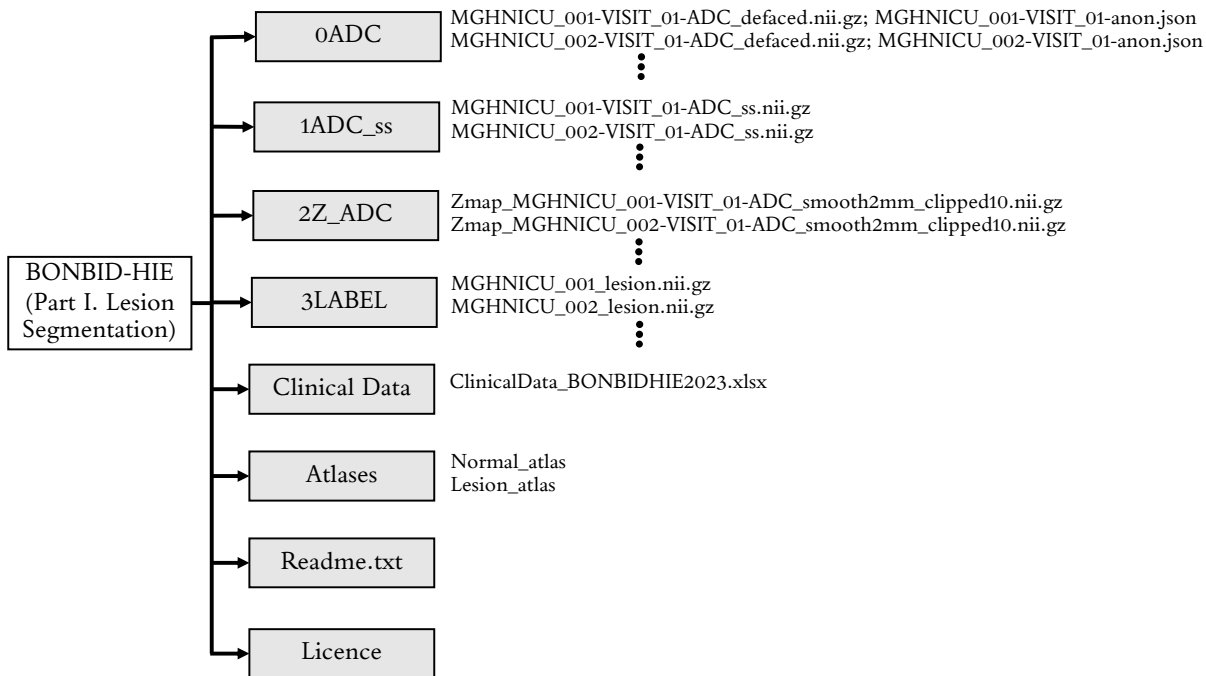


Figure 6. Folder structure of the BONBID-HIE dataset (Part I. Lesion Segmentation).

145 Technical Validation

146 Representativeness of patient cohort

147 Our data is representative of HIE cohorts in the developed countries. At least three characteristics of our data agree with
 148 documented clinical knowledge about HIE.

149 **Lesion distribution in space agrees with clinical knowledge.** Our statistical lesion atlas in Figure 4 shows that HIE
 150 lesions can occur anywhere in the brain. The regions most frequently injured here included the basal ganglia, internal capsules,
 151 thalamus, temporal lobes, cerebral white matter, brainstem, and cerebellum (red, orange, and yellow regions in Figure 4).
 152 This lesion atlas map coincides with clinical knowledge of brain regions often vulnerable to HIE injuries [2, 52, 6]. Indeed,
 153 HIE-related injuries in these regions have been key criteria in expert MRI scoring systems, which are used to assess the severity
 154 of HIE. Examples include the NICHD Neonatal Research Network (NRN) [2], the Barkovich [53], the Weeke-deVeries [3], and
 155 the Trivedi [54] scoring systems. In addition, lesions appeared in less than 35% of the patients at any given voxel, according to
 156 the color bar in this figure. This confirms the clinical knowledge that HIE lesions are diffuse, spatially distributed, and almost
 157 half to two-thirds of the HIE patients are mild to moderate, at least in patients in the USA [50, 51].

158 **Lesion distribution in time agrees with clinical knowledge.** Figure 7(a) shows the percentage of the whole brain volume
 159 being lesioned at different postnatal ages. The lesion percentage in the ADC maps came down to almost 0 in the 9 patients
 160 who underwent MRI scans after postnatal day 7. This agrees with the clinical knowledge that HIE-related lesions are more
 161 detectable in ADC during 0-7 postnatal days or in T1/T2-weighted images than in relatively later scans (after postnatal day
 162 7) [55, 51, 2].

163 **ADC evolution with age agrees with clinical knowledge.** Figure 7(b) shows the whole-brain average ADC values of all
 164 patients (each dot is a patient). In normal cohorts, ADC values drop rapidly in the early postnatal life (see Figures 4 and 5
 165 in [36], and Figures 3 and 4 in [46]). However, the presence of HIE-related abnormalities disrupted this trend – HIE patients
 166 undergoing earlier MRIs (0-7 postnatal days) had decreased ADC values in a larger percentage of the brain (Figure 7(a)), so the
 167 ADC values in 0-7 postnatal days were at similar or even lower levels than ADC values in 7-14 postnatal days among HIE
 168 patients (Figure 7(b)). This has also been documented in HIE literature [56, 57].

169 Utility of ADC maps and Z_{ADC} maps

170 To demonstrate the utility of the computed Z_{ADC} maps, we compared the accuracies of using ADC or Z_{ADC} maps for lesion
 171 segmentation. We attempted simple thresholding of ADC and Z_{ADC} maps, at several threshold values, for segmenting HIE
 172 lesions. Although simple, thresholding-based segmentation accuracy is a strong indicator for segmentation accuracies in more

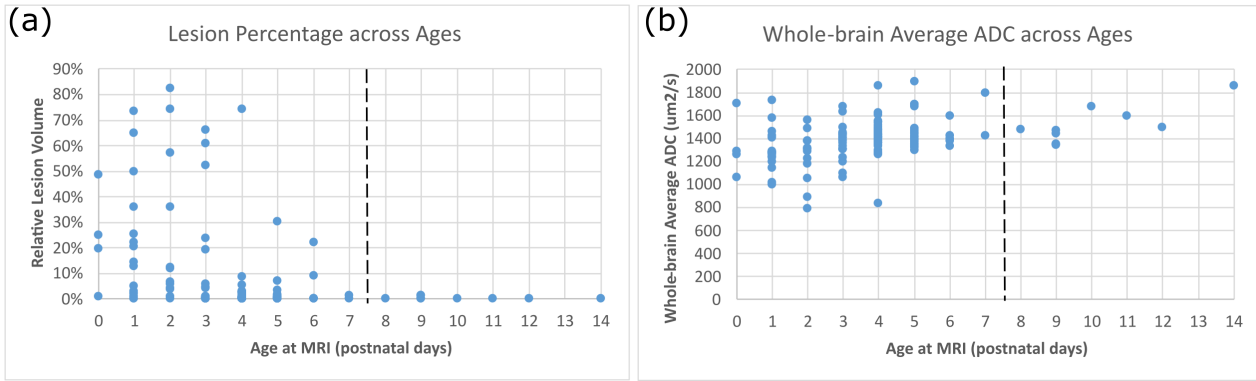


Figure 7. Representativeness of our cohort for (a) lesion distribution across ages; and (b) whole-brain ADC values across ages. In both panels, each dot denotes a patient in our cohort.

173 sophisticated machine/deep learning algorithms [58]. For ADC maps, we used different thresholds ranging from 800-1100
 174 $\mu\text{m}^2/\text{s}$, as suggested in the literature [55, 59, 60, 61, 62]. For Z_{ADC} maps, we used thresholds -1.5, -2, and -2.5. Voxels in
 175 patient MRIs with ADC values 1.5 to 2.5 standard deviations below the average ADC values from healthy controls were
 176 considered abnormally low and hence, lesioned. The choice of thresholds around -2 in Z_{ADC} maps was also based on the normal
 177 distribution of ADC values at each voxel across subjects [36, 40].

178 We evaluated the accuracy of these maps compared to expert-annotated ground-truth masks using the Dice coefficient,
 179 sensitivity, and specificity. Results are shown in Figure 8. Here, the gray boxplots are the accuracy measurements when ADC
 180 maps were thresholded between 800 and 1100 $\mu\text{m}^2/\text{s}$ (50 $\mu\text{m}^2/\text{s}$ intervals). The blue boxplots are the accuracy measurements
 181 when the Z_{ADC} maps were thresholded at -1.5, -2, and -2.5. Figure 8 demonstrates: (i) both ADC and Z_{ADC} have value in helping
 182 segment the HIE-related lesions, since the specificity from simple naive thresholding-based segmentations was comparable to
 183 those from machine learning-based algorithms [20], although the Dice and sensitivity were lower; (ii) Z_{ADC} maps thresholded
 184 at -2, the most intuitive and straightforward threshold value, yielded the highest Dice (0.54 ± 0.28), followed by Z_{ADC} maps
 185 thresholded at -2.5 (Dice 0.39 ± 0.25); (iii) Z_{ADC} maps thresholded at -1.5 yielded the highest $\frac{\text{sensitivity}^2 + \text{specificity}^2}{2}$ (0.82 ± 0.02)
 186 compared with any ADC thresholds, followed by Z_{ADC} maps thresholded at -2 (0.76 ± 0.03); and (iv) overall, across all
 187 thresholds, Z_{ADC} maps showed a higher area under the curve (AUC: 0.936). This shows that Z_{ADC} maps – anatomy-normalized
 188 ADC images – carry the potential to improve lesion detection accuracy.

189 Code Availability

190 Data is available as part of the 1st BONBID-HIE MICCAI challenge (<https://hiechallenge.github.io/>), to be
 191 held in October 2023. For the training sub-cohort, all MRI (raw and derived) and expert lesions annotations are available. For
 192 the testing sub-cohort, only the MRI (raw and derived) data is available at the time this manuscript is drafted, and the expert
 193 lesion annotations will be made available after the challenge in October 2023.

194 Codes to automatically calculate the evaluation metrics are available in BONBID-HIE 2023 MICCAI challenge (<https://bonbid-hie2023.grand-challenge.org/>). The repository in the challenge contains scripts to read the images,
 195 visualize them, and quantify the algorithm's performance with the same metrics used in the challenge to rank participants.
 196

197 Acknowledgements

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199 Competing interests

200 None.

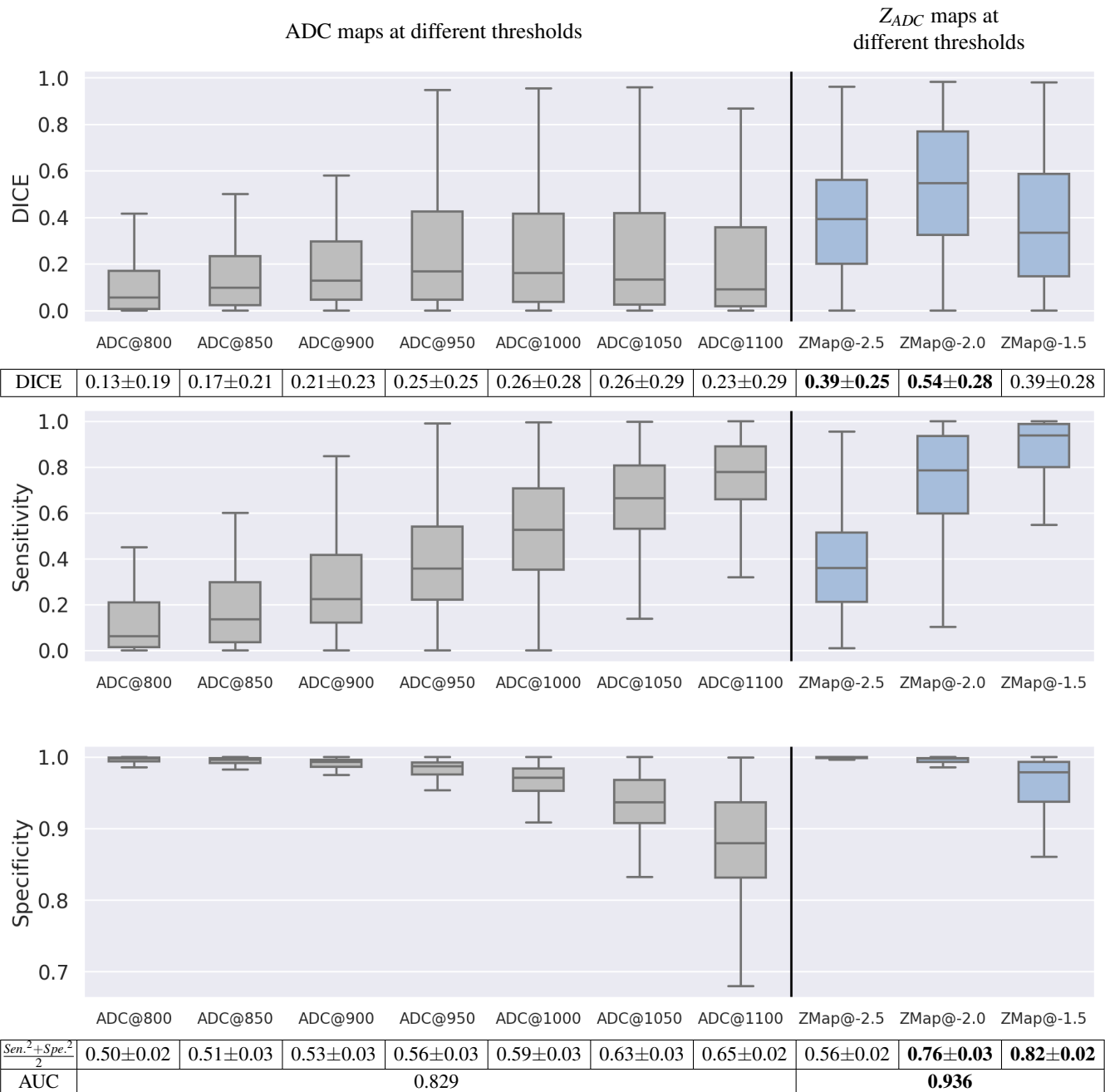


Figure 8. Accuracy of thresholding-based lesion segmentation on ADC and Z_{ADC} maps with different threshold values. Bold texts in the tables beneath the figure panels highlight the two scenarios with the highest Dice scores, the two scenarios with the highest and most balanced sensitivity and specificity metrics, and the parametric map (ADC or Z_{ADC}) with the highest area under the receiver-operating-characteristic curve (AUC).

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