



Data Resource Profile

Data Resource Profile: Better Outcomes Registry & Network (BORN) Ontario

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Data resource basics

The Better Outcomes Registry & Network (BORN) Ontario collects pregnancy, birth and newborn information and outcomes data for nearly all births in Ontario, Canada. With nearly complete capture of the approximate 140 000 births each year in Ontario—~40% of births in Canada—BORN Ontario is the largest perinatal registry in the country.¹ The origins of BORN Ontario date to the 1980s as a system for tracking regional variation in pregnancy and childbirth outcomes across Eastern Ontario. BORN Ontario was originally named the Niday Perinatal Database in honour of its founder, Dr Patricia Niday, who, as director of one of Ontario's regional perinatal programmes, advocated tirelessly for its provincial adoption. The regional programme gradually expanded data capture and launched the Ontario Perinatal Surveillance System (OPSS). By 2006, most of the province was contributing data to the OPSS for analysis and reporting.²

In 2009, with funding from the Ontario Ministry of Health and Long-Term Care, OPSS was designated a prescribed registry under the province's Personal Health Information Protection Act, and rebranded as BORN Ontario. Prescribed registry status granted BORN Ontario the authorization to collect, use and disclose personal health information without consent to facilitate and improve healthcare. The new organization amalgamated the data systems of four programmes servicing the maternalnewborn population: the Prenatal Screening Program, the Fetal Alert Network (high-risk antenatal healthcare for pregnancies complicated by congenital anomalies), the

Key Features

- The Better Outcomes Registry & Network (BORN) Ontario was developed to collect pregnancy, birth and newborn information and outcomes data to facilitate and improve perinatal care in Ontario, Canada. With near-complete capture of all births in Ontario, it has evolved to be an invaluable resource for performance measurement, quality improvement, surveillance and research.
- The BORN Information System was launched in 2012 as the amalgamation of data systems from provincial programmes servicing the maternal-newborn population. BORN Ontario also maintains the legacy data sets of its predecessors (2006–2012) and other data holdings have since been added.
- Data are submitted regularly by >250 hospitals, fertility clinics, birth centres, midwifery practice groups, primary care
 organizations, and prenatal- and newborn-screening laboratory and treatment centres. Over 1.3 million maternal
 infant records have been captured to date.
- Data include maternal demographics, health behaviours, in vitro fertilization assistance, prenatal screening, pregnancy interventions and complications, intrapartum events, peripartum outcomes, intensive-care and newborn-screening information. Record-level data can be linked to a wide range of provincial administrative and clinical data sets to track individuals from conception to birth and across the lifespan.
- Aggregate and record-level data are available upon request via the Data Access Request Team. BORN Ontario data
 dictionaries are openly available to browse online.

Ontario Midwifery Program and the Niday Perinatal Database. These programmes worked together to normalize their data and build a robust, web-based data-collection and reporting system. In April 2012, the BORN Information System was launched to collect, manage, protect and share critical data about every pregnancy, birth and child in Ontario. Newborn screening test results from Newborn Screening Ontario were also included at launch, representing the first time such information was linked to perinatal data sets in Ontario. A reporting portal for data contributors went live in 2013.

BORN Ontario has evolved to be a secure data platform with multiple data holdings that can be linked to connect individual health records from conception to birth and into early childhood. The Canadian Assisted Reproductive Technologies Registry (CARTR), which collects data on *in vitro* fertilization (IVF) cycles and birth-outcomes data for fertility clinics across Canada, was added in January 2013 and rebranded CARTR Plus. Other data holdings have since been added, including primary care data on young children and results of standardized screening and assessments of child development through the provincial Healthy Babies Healthy Children programme. BORN Ontario also maintains legacy data sets from the founding five partner programmes (last data entry 31 March 2012) and CARTR (January 2001— December 2012).

The present-day BORN Information System collects data from >250 hospitals, fertility clinics, birth centres, midwifery practice groups, primary-care organizations, and prenatal- and newborn-screening laboratory and treatment centres spanning all levels of care from pre-pregnancy to early childhood. To date, >1.3 million maternal-infant records have been captured. The following sections will focus on the BORN Information System with reference to the Niday Perinatal Database for historical context.

Data collected

BORN data are collected on a voluntary basis from healthinformation custodians involved in the care of children, newborns and their mothers. Data are submitted through several mechanisms including manual data entry into a reporting portal by staff in birthing units and midwifery practice groups, Health Level Seven (HL7) data feeds, and automated extraction and uploads from electronic health record systems, where available. Data are submitted in close to real time, or uploaded in batches, and include maternal demographics, health behaviours, any IVF assistance, prenatal screening, pregnancy interventions and complications, intrapartum events, peripartum outcomes, neonatal intensive care unit (NICU) care and newborn-screening data, each of which is classified by the nature of the clinical encounter with the healthcare system (Figure 1). Linkage of data elements and encounters across data sources is facilitated by a robust linking and matching algorithm utilizing unique individual (mother and child), pregnancy and birth identifiers that are assigned upon the first record entry into the BORN Information System. Some data elements are common across encounters, permitting the pre-population of some data fields to reduce duplication of data-entry



Figure 1 Better Outcomes Registry & Network (BORN) Ontario data sources and architecture. BORN Ontario data are submitted on a voluntary basis by health-information custodians from across Ontario servicing the maternal–child population. Data are classified by the nature of the healthcare encounter and are linked to create aggregate records for each woman and child. CARTR, Canadian Assisted Reproductive Technologies Registry; MFM, maternal fetal medicine; NICU, neonatal intensive care unit; NIPT, non-invasive prenatal testing; SCN, special care nursery; HBHC, Healthy Babies Healthy Children.

efforts. This model enables both programme-specific analyses by encounter and analysis of aggregated data by population and health system. A description of the data available in BORN is provided in Table 1.

Data quality

BORN Ontario data are used to support clinical programme management, benchmarking, evaluation and quality improvement across the province of Ontario. The data may also to be used for research. As such, BORN Ontario upholds a comprehensive framework to promote quality in all aspects of data collection, analysis, use and disclosure of information.⁸ Five essential dimensions of data quality are upheld (Figure 2).

Audits of the Niday Perinatal Database and BORN Information System have demonstrated good agreement with patient charts and clinical administrative hospital databases.^{9–12} An external audit by Public Health Ontario concluded that the quality of BORN data exceeded that of other sources of reproductive-health information.¹² A reabstraction study of 29 variables from records submitted in 2014–2015 found >90% agreement between BORN records and patient charts for over three-quarters of the audited variables.¹¹ The remaining variables demonstrated fair to moderate agreement. These findings have informed ongoing improvements to BORN data dictionaries, dataentry guidelines and quality-control processes.

The BORN Information System applies a series of validation rules to ensure data quality. Logic rules assess the compliance of data elements collected within a given healthcare encounter. Conformance rules ensure that mandatory elements are completed. Parameter rules compare entered values against reference tables to ensure that they are within acceptable ranges before they are submitted. The system also applies aggregation rules to determine how data elements across multiple encounters (e.g. prenatal screening, general antenatal visits and delivery) are consolidated into a single birth record. Where there is discordance in data elements that are recorded in more than one encounter, aggregation rules provide guidance on which values take precedence. Thus, having some data elements captured multiple times across the pregnancy and birth continuum also helps to increase data accuracy and completeness.

Support mechanisms are in place for submitting organizations to identify and resolve errors in their data entries. The data-validation rules described above facilitate the generation of reports and flags for records identified with missing encounters, possible errors or different data entered for the same element in separate encounters so that errors can be reconciled. All data must be formally acknowledged by the submitting organization and, each month, submitters review their records to ensure that there are no outstanding errors. As a final checkpoint in this process, BORN data analysts conduct additional data checks at the end of each fiscal year.

Finally, online help tips and data-element definitions are available within the data-entry system and regional coordinators are available to answer questions related to data entry, reconciliation and acknowledgement processes, and the implementation of system enhancements. Regional

Table 1 Summary of data collected by the Better Outcomes Registry & Network (BORN) information system

Encounter type or data holding	Description		
Assisted reproductive technology (ART) ^{3,4}	Collects information on <i>in vitro</i> fertilization treatment cycles from fertility clinics across Canada col- lected into the Canadian Assisted Reproductive Technology Register (CARTR Plus)		
	Also includes record-level details on any ART treatment cycles from Ontario fertility clinics that result in a pregnancy that is delivered at >20 weeks' gestation that are automatically linked with other records within the BORN Information System		
Antenatal general	Captures information relevant to the antenatal period (i.e. obstetrical history, exposures, complica- tions, etc.)		
Antenatal specialty	Captures maternal, pregnancy and fetal information for women who require specialized care manage ment at a high-risk maternal fetal medicine or prenatal genetic clinic		
Prenatal screening and follow- up ^{5,6}	Prenatal screening records for every pregnant individual who receives any form of prenatal screening in Ontario		
	Includes information from laboratory and ultrasound testing, clinical assessment, genetic counselling and diagnostic testing from prenatal screening, as well as screening and diagnostic results		
Non-invasive prenatal testing ⁷	Information captured during non-invasive prenatal testing for fetal genetic disorders		
Cytogenetics	Information related to the clinical indications and results of maternal and child cytogenetic testing		
Labour and birth	Captures information relevant to the pregnant person's spontaneous or induced labour and birth ex- perience through to the first hour postpartum whether this occurs in hospital, at home or in a birth centre		
	Midwifery-client specific data are also captured in the midwifery encounter		
Birth–child	Documents a child's birth experience through to the first hour postpartum		
Postpartum mother	Documents the course of care experienced by the mother from 1 hour post birth to discharge from hospital		
Postpartum child	Documents the course of care experienced by the child from 1 hour post birth to discharge from hospital		
Newborn screening and diagnostic evaluations	Screening tests and results from the provincial newborn-screening laboratory (including hearing and congenital heart disease)		
	Information on short-term follow-up, clinical status, diagnostic information and treatment plans for infants who received a positive newborn screen are also available		
Neonatal intensive care/special care units	Information collected during infant stays in neonatal intensive care units and special care nurseries		
Midwifery practice group data	Data on prenatal and birth care (home, hospital and birth-centre births) provided by midwives in Ontario		
Well Baby Well Child visit	Information obtained through regular assessments of a baby or child's development		
information	Includes age, height, weight, vaccinations and documentation of developmental milestones using the standardized measures (Rourke Baby Record Ontario, Nipissing District Developmental Screen)		
Healthy Babies Healthy Children (HBHC) programme	Information collected from completion of the standardized HBHC Screening Tool, which facilitates identification of risk factors that may affect a child's healthy development and referral to community programmes and services		
	Screening is voluntary and consent-based		
	BORN facilitates the transfer of this information from hospitals/homes to public-health units who provide the follow-up		

Citations are provided where data elements have been previously validated or described in detail.

coordinators are distributed throughout the province and work directly and regularly with the data submitters and the regional perinatal networks using the data to improve care. A 24-hour Helpdesk is also available.

Data privacy and security

BORN Ontario is a prescribed registry established under the Ontario Personal Health Information Protection Act, 2004 A.¹³ As such, personal health information within the

BORN network is protected by administrative, physical and technological controls that adhere to industry best practices for privacy and security. BORN Ontario has implemented a rigorous programme to protect personal health information in its custody from theft, loss, unauthorized access, copying, modification, use, disclosure and disposal. Regular audits and investigations are conducted to monitor and manage privacy compliance. BORN Ontario's information practices and privacy policies are assessed by the Information and Privacy Commissioner of Ontario every 3 years.

Timeliness	How current or up-to-date data is at the time of release and whether the data is available for user needs within a reasonable time-period.	
Accuracy	How well information within, or derived from the database, reflects the reality it was designed to measure.	
Comparability	The extent to which data are consistent over time and entered using standard conventions making it comparable to other databases.	
Usability	The ease with which the data is understood and accessed.	
Relevance	The degree to which the data meets the current and potential future needs of users.	

Figure 2 Better Outcomes Registry & Network (BORN) Ontario dataquality framework. The data-quality framework is based on five dimensions to ensure that user decision-making is based on current, valid, reliable and relevant data.

Governance and funding

The Children's Hospital of Eastern Ontario (CHEO) provides the formal governance structure and support for BORN Ontario operations. The BORN Ontario Executive Team works in cooperation with leadership at CHEO to provide medical, scientific and administrative oversight of day-to-day operations. Core funding is provided by the Ontario Ministry of Health and Long-Term Care.

Data resource use

Contributing hospitals, laboratories, clinics and birth centres have access to their own data within the BORN Information System. These data are used locally for practice audits and to support quality-improvement initiatives. In addition, various organizations and external partners may use BORN Ontario data for planning and management of the healthcare-delivery system at organizational, provincial or other regional levels. These efforts have resulted in numerous successful initiatives to enhance the quality of maternal-newborn care delivery in Ontario through data-quality assessments,⁹⁻¹¹ the introduction of new health services and programmes¹⁴⁻¹⁷ and technology-based interventions.¹⁸⁻²² The BORN Ontario registry is also an invaluable research resource for addressing maternal and child health issues. BORN data have been used extensively to address research topics including the developmental origins of health and disease, complications of pregnancy and childbirth, and evaluating short- and long-term maternal and child health outcomes.

BORN dashboards

BORN has developed two electronic audit and feedback systems that provide hospitals offering maternal, newborn and NICU care with site-specific feedback on a near real-time basis to facilitate practice change on key performance indicators.¹⁸⁻²¹ The BORN Maternal Newborn Dashboard was launched in 2012 with six key performance indicators and has since become an instrumental tool for decreasing variation in clinical practice and improving patient outcomes across the province. Evaluation of the Maternal Newborn Dashboard 2.5 years after its implementation demonstrated corresponding improvements across several key performance indicators. These include: rates of episiotomy, post-date induction at <41 weeks, repeat caesarean section in low-risk women before 39 weeks and appropriately timed group B streptococcus screening.¹⁸ A NICU Dashboard was also launched in 2018 with two key performance indicators to support clinical-practice improvement for neonatal care. Each dashboard displays a hospital's performance on each key performance indicator relative to provincial benchmarks, provincial-level rates as well as peer-performance data from hospitals providing similar levels of maternal or neonatal care and/or of similar birth volume. Engagement with local health professionals and organizations is ongoing and helps to ensure uptake of these dashboards and meaningful practice change.^{19,20}

Perinatal research

Since its inception, BORN data have been used in Canada and internationally to produce nearly 100 high-impact publications with broad medical and scientific reach. These publications cover a diverse array of topics including, but not limited to, prenatal screening and diagnostic testing^{7,23}; breastfeeding and influencing factors^{24,25}; use and outcomes of assisted reproduction^{26–30}; trends in obstetrical interventions and factors influencing use^{31–33}; and factors influencing maternal and neonatal health outcomes such as environmental exposures,^{34–36} influenza immunization during pregnancy,^{37–39} substance use,^{40,41} hypertension,^{42,43} diabetes^{44–47} and obesity/gestational weight gain.^{48–50} A compendium of published research using BORN data is available online (https://bit.ly/2UXEX3I).

Data linkage: long-term follow-up of women and infants

BORN data can be linked with other provincial data resources to facilitate long-term follow-up of women and their children. ICES (www.ices.ca) is a provincial data repository containing longitudinal, coded and linkable clinical and administrative records encompassing nearly all interactions that Ontarians have with the publicly funded healthcare system. Population-based health surveys, research-specific and registry data are also integrated, permitting evaluation of health policy and care delivery as a whole across the province. In 2014, the Niday historical data set (2006/07-2011/12) was transferred to ICES. A 2-year transfer of BORN Information System data spanning 2012/13-2013/14 fiscal years was completed in early 2015; a second data transfer (2014/15-2018/19) took place in 2020. The new data are scheduled for release in 2021. Deterministic and probabilistic linkage of individual maternal and newborn records to master registration files at ICES facilitates linkage of BORN data to all other data sets held by ICES. Thus, linkage of these two repositories provides a unique opportunity to study population-level maternal and infant health outcomes ranging from the fertility treatment and perinatal period to many years after birth. Recent applications of ICES-BORN data include examining the effect of frequent blood donations before pregnancy on maternal and newborn outcomes⁵¹; evaluating the association between cannabis exposure in pregnancy and neurodevelopmental outcomes in children⁵²; assessing the 5-year health outcomes of children born to mothers who received the 2009 pandemic H1N1 influenza vaccination during pregnancy³⁸; exploring the link between prenatal and earlylife exposures to ambient air pollutants and the development childhood asthma⁵³; the association between intrapartum epidurals and postpartum depression⁵⁴; and risk of breast cancer following thoracic computed tomography in the perinatal period.⁵⁵ The potential applications of these linked data include interrupted time-series analyses and using BORN to collect data for clinical trials and other interventions. Given that BORN Ontario is a relatively new entity, the potential use of these linked data will only continue to grow, particularly as other provincial data holdings become available.

Strengths and weaknesses

Ontario, Canada has rich network of health-data repositories that can be used to provide comprehensive insight into population health and healthcare delivery. The BORN Ontario registry is a component of the network, providing the capacity to link pregnancy and newborn health records across multiple health systems and track individuals from conception and birth across their lifespans. The breadth of data held by the BORN Ontario registry makes it a foundational resource for answering perinatal health-related questions. Strengths include: capture of all hospital births in Ontario (>99% of all births in the province) and all midwifery-attended home and birth-centre births; capture of extensive information about maternal and newborn characteristics, health histories and behaviours, outcomes and health-services utilization; integration of multiple data holdings spanning the continuum of maternal-child healthcare delivery; and a nearly 10-year data-collection time frame. Notably, BORN data have been validated and demonstrate consistency with other clinical administrative data sets.

The BORN Ontario birth registry is not without limitations. BORN elements are subject to incomplete or duplicate data entries, misclassification and under-reporting. The BORN Registry is reliant on data sourced from medical records, clinical forms and patient interviews, and, if data are not collected or entered into a patient's records, they will not be available for abstraction into the BORN Information System. In addition, where manual entry may be performed by clerk staff or nurses, variability in the interpretation of case definitions can affect data quality, and incomplete or duplicate data entry and misclassification are a reality. Data entry errors and duplicate data entries are cleaned when identified. Collection of pre-pregnancy body mass index is limited by early-pregnancy weights or patient self-report, although recent efforts to standardize how this information is collected have improved the data quality for this variable. Information on miscarriage and therapeutic abortions is historically difficult to ascertain, and under-reporting of socially sensitive variables including those related to substance use/abuse, intimate-partner violence and sexually transmitted infections is also likely. Last, socio-economic information at the individual or family level and healthcare-provider identifiers are not available in BORN. Area-level socio-economic information (e.g. education, income, marginalization and deprivation indices) can be obtained, however, through linkage to other resources including Canadian census data.

Data resource access

Access to legacy data sets and the BORN Information System

Data are accessible to contributors through the BORN Information System reporting portal. Each contributor has a designated lead who can grant role-based access for members of their group. Those who do not have access to the reporting portal or who require data that are not available through this mechanism may pursue alternative processes to access BORN Ontario data resources. Legacy data sets and BORN Information System data are available to external requestors to support research, programme evaluation, policy development and surveillance (Table 2). All requests to BORN Ontario are managed in accordance with provincial privacy requirements.

Data requests can be made to the BORN Data Analysis & Request Team and initiated through submission of a data request inquiry form (https://forms.bornontario.ca/Data-

	Legacy data holdings	BORN Information System
Description	 Includes record-level data from: Niday Perinatal Database the Ontario Midwifery Program Prenatal Screening Ontario and historical non-invasive prenatal-testing and cytogenetic-testing results the Canadian Assisted Reproductive Technologies Register (CARTR) 	Includes record-level data from birthing hospitals, midwifery practice groups, birthing centres, fer- tility clinics, prenatal- and newborn-screening laboratories, follow-up clinics, clinical pro- grammes and primary-care organizations
Time frame Data dictionary	April 2006–March 2012 https://bit.ly/2whlSRu	April 2012–present https://bit.ly/2155jec

Table 2 Available data sets at Better	Outcomes Registry &	Network (BORN) Ontario
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Inquiry-Form). From here, the application process varies, depending on the type of information requested (aggregate tables or record-level data) and the support required (Figure 3). BORN coordinators and data analysts work closely with external requestors to navigate data access approval processes, prepare analytical data sets and conduct analyses. At present, BORN Ontario does not provide a platform through which external requestors can interact with the data for research purposes. Research data sets are securely and electronically transferred to external requestors following completion of all of the necessary approval processes.

Data dictionaries for legacy data elements are available for download. The BORN Information System data dictionary is available online as an interactive library and may also be downloaded in portable document form. A data dictionary improvement project is currently underway. Notably, the data dictionary and data elements available from the legacy data sets and the BORN Information System differ. New data elements and updated terminology were added to the BORN Information System in 2012 and may not be available in the historical data. It is not always possible to link some of the older data elements to the new BORN Information System.

Access to data relating to Indigenous peoples in Canada

Data on pregnant individuals and newborns with a postal code where the majority of the population identifies with an Indigenous Nation (First Nation, Inuit and Métis peoples) are only available for public-health activities or research analysis when there is evidence of stakeholder engagement and consent of the Indigenous Nation(s). BORN Ontario has a responsibility to ensure that the information it collects is managed in a culturally appropriate and respectful manner, and thus adheres to the principles of ownership, control, access and possession (OCAP[®]) and other similar Indigenous frameworks for conducting research pertaining to Indigenous people. The decision to suppress Indigenous data in BORN is based on guidance from Indigenous stakeholders on the collection, interpretation and governance of data related to their community members and communities as a whole.⁵⁶ Resources are dedicated to facilitating data release to Indigenous stakeholders for programme planning and delivery of maternal, newborn and early-childhood services and does not otherwise release information without evidence of engagement and consent by a community stakeholder.

Data access fees

Costs associated with preparing aggregate data and record-level data sets for external requestors depend on the nature and complexity of the request. Following consultation and an assessment of the project needs, cost estimates can be provided for research-funding applications, for funded research projects and other quality initiative and policy work. Data access fees include consultation, risk assessment and de-identification, data analyst time for file preparation and verification, and an administration fee.

Summary

BORN Ontario is a primary and authoritative resource for reproductive health information and data in Ontario, Canada. The BORN Information System is an accurate and trusted tool for evaluating care delivery and health outcomes of mothers and their newborns. It can further be used to support data-driven change-management strategies in perinatal healthcare policy and programming. Importantly, BORN Ontario data are accessible to local,



Figure 3 Accessing Better Outcomes Registry & Network (BORN) Ontario data. External parties may submit requests for aggregate or record-level data. Requests can be initiated through submission of an online form. Turnaround times and data access fees may vary, depending on the nature and complexity of the request. Requests to access record-level data from BORN Ontario data holdings may be subject to additional approvals.

national and international investigators looking to address research questions relevant to maternal, newborn and paediatric populations.

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Conflict of interest

None declared.

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