**TITLE PAGE**

**Title:** Health Outcomes and drug utilisation in children with Noonan syndrome: a European cohort study

**Authors:**

Michele Santoro1, Ingeborg Barisic,2, Alessio Coi1, Joachim Tan3,4, Ester Garne5, Maria Loane6, Ljubica Odak7, Maria Valentina Abate1,Elisa Ballardini8,Clara Cavero-Carbonell9, Miriam Gatt10, Mika Gissler11,12,13, Kari Klungsøyr14,15,Nathalie Lelong16, David Tucker17, Diana Wellesley18, Joan K. Morris4

1Unit of Epidemiology of Rare diseases and Congenital anomalies, Institute of Clinical Physiology, National Research Council, Pisa, Italy;

2Children's Hospital Zagreb, Centre of Excellence for Reproductive and Regenerative Medicine, Medical School University of Zagreb, Zagreb, Croatia;

3NIHR Great Ormond Street Hospital Biomedical Research Centre, UCL GOS Institute of Child Health, London, UK;

4School of Health and Medical Sciences, City St George’s University of London, London, UK;

5 Department of Paediatrics and Adolescent Medicine, Lillebaelt Hospital, University Hospital of Southern Denmark, Kolding, Denmark.

6Faculty of Life and Health Sciences, Ulster University, Northern Ireland, UK;

7Department of Medical and Laboratory Genetics, Endocrinology and Diabetology with daily care unit, Children's Hospital Zagreb, Zagreb, Croatia;

8Neonatal Intensive Care Unit, University Hospital of Ferrara IMER Registry (Emilia Romagna Registry of Birth Defects) Department of Medical Sciences, University of Ferrara, Italy;

9Rare Diseases Research Unit, Foundation for the Promotion of Health and Biomedical Research in the Valencian Region, Valencia, Spain;

10Directorate for Health Information and Research, G’Mangia, Malta;

11THL Finnish Institute for Health and Welfare, Department of Knowledge Brokers, Helsinki, Finland;

12Region Stockholm, Academic Primary Health Care Centre, Stockholm, Sweden;

13Karolinska Institutet, Department of Molecular Medicine and Surgery, Stockholm, Sweden; 14Department of Global Public Health and Primary Care, University of Bergen, Norway;

15Division of Mental and Physical Health, Norwegian Institute of Public Health, Bergen, Norway;

16Université de Paris, CRESS-Epopé, INSERM, INRA, Paris, France;

17Public Health Wales, Swansea, United Kingdom;

18University Hospital Southampton, Wessex Clinical Genetics Service, Southampton SO16 5YA, UK;

**Corresponding author**: Michele Santoro, Institute of Clinical Physiology, National Research Council, Via Moruzzi 1[, 56124 Pisa (Italy),](mailto:asmith@example.com) michele.santoro@cnr.it, phone number: 0039503158120.

**Keywords:**

Noonan syndrome; cohort; survival; hospitalization; surgeries; prescriptions

**ABSTRACT**

**Background.**Noonan Syndrome (NS) is a rare multisystemic disorder with heterogeneous phenotypic manifestations. The aim of this study was to analyse rates of survival, hospitalisation, surgeries and prescriptions in children born with NS in the first 10 years of life.

**Methods.** This is a multi-centre population-based cohort study. Data on175liveborn children diagnosed with NS from 11 EUROCAT congenital anomaly registries were linked to healthcare databases. Each registry applied a common data model to standardise data and run common syntax scripts to produce aggregated results which were pooled using random effects meta-analyses.

**Results.** Mortality rates were high in the first year of life with 5.4% (95%CI 1.5%-10.1%) of children dying before the age of 1 year with a further 2% dying up to age 5. In the first year, 87.9% (95%CI 75.3%-94.3%) of children were hospitalized and the median Length Of hospital Stay (LOS) was 15.3 days (95%CI 9.3-21.2). After the first year, the proportion of children hospitalized remained higher than 70%, but the LOS decreased to 1.3 days per year. In the first 5 years, 65.2% of children underwent a median of two surgical procedures. The median age at first surgery was 29 weeks. The proportion of children with an antibiotic prescription increased from 53.6% at age 1 to 62.4% yearly until 4 years of age.

**Conclusions**. Children with NS have high mortality and morbidity not only in the first year of life but also up to five years of age. This study evaluated the health burden of NS and provided information for clinicians, health-care providers and families.

**MANUSCRIPT**

**BACKGROUND**

Noonan Syndrome (NS) (MIM # 605275) is typically an autosomal dominant multisystem disorder with heterogeneous phenotypic manifestations which represent a major burden for affected patients and their families.1 NS belongs to the family of RASopathies, a group of conditions caused by gain-of-function pathogenic variants in genes encoding components or regulators of the RAS/mitogen-activated protein kinase signalling pathway.2,3 Clinical characteristics may include distinctive facial features, broad or webbed neck, congenital heart defects, chest deformity, renal anomalies, cryptorchidism, lymphatic malformations, bleeding diathesis, short stature and developmental delay/learning difficulties. The phenotype is variable and mildly affected individuals may not be diagnosed until adulthood. Diagnosis is typically based on clinical features, but molecular genetic testing can confirm the diagnosis in about 80% of cases.4-10 It is necessary that different paediatric subspecialties are familiar with the health burden and course of NS to ensure appropriate parental counselling, multidisciplinary monitoring and treatment.

Children and adults with NS need to receive regular medical care and follow-up to monitor and manage their health.6,7,10 Medications may be prescribed, but specific information on prescriptions issued to individuals with NS is lacking.12-15

The mortality rate can vary depending on several factors, including the severity of cardiac abnormalities that are present in 50 to 80% of individuals.9,16 According to a study that followed a large cohort of individuals with RASopathies, the overall mortality is relatively low.17 However, certain cardiac complications, such as Hypertrophic CardioMyopathy (HCM), can increase the risk of mortality.17,18 A report on patients with NS and HCM showed a significantly worse risk-adjusted late survival rate compared to individuals with nonsyndromic HCM.19 Additionally, a study analysing outcomes in children with NS and HCM found that patients with NS and HCM had a worse risk profile at presentation, resulting in significant early mortality.20 Additional population-based studies are needed to further understand the mortality risk and long-term outcomes in population-based cohorts of newborns with NS.

Hospitalisation may be necessary for those who experience complications related to their condition, such as congenital heart defects, feeding problems and failure to thrive in the first year of life. Some individuals with NS may require surgical procedures.16,21 It is suggested that NS is associated with increased hospital stays for paediatric cardiac surgery which has implications for costs. One study found that NS accounted for 1.6% of paediatric cardiac surgery admissions, and patients with NS tended to have longer hospital stays, higher total charges, and increased inpatient mortality compared to those without NS.22 However, there is still little evidence to provide specific information based on a population-based approach about hospitalisation and surgery rates in general for all individuals with NS with or without congenital heart defects.

The study presented here is part of the EUROlinkCAT project that aimed to investigate the health outcomes of European children born with major Congenital Anomalies (CA) by linking data from population-based registries to electronic health care databases.23

The aim of this study was to analyse rates of survival, hospitalisation, surgeries and prescriptions in children born with NS in the first 10 years of life.

**METHODS**

This is a population-based data-linkage cohort study. Data were available on children born with NS (ICD10 BPA code Q87.14 and ICD9 BPA code 759.896) in the period 1995-2014 from 11 European Surveillance of Congenital Anomalies (EUROCAT) registries24,25 in 7 European countries. EUROCAT registries collect standardised data on all major CA cases diagnosed by the first year of age in their population using multiple sources of ascertainment according to EUROCAT guidelines.26

Data from participating registries were linked to data in local healthcare databases covering the registry's geographical area (mortality, hospital admission and discharge data, prescription data). Linked data on outcomes of interest were included up to the child’s 10th birthday or to 31st December 2015 (whichever was earlier). A detailed description of the linkage methods used in the EUROlinkCAT project has been published elsewhere.23,27,28 Only years with good quality healthcare data and a high proportion of successful data linkage were included in the study period and the overall successful linkage was higher than 95%.27

For the investigation of survival, data on mortality were obtained through electronic linkage with vital statistics and mortality databases used in the regions and countries covered by the registries. A detailed description of the methods has been published elsewhere.29-31

For the analysis of hospitalisation and surgical procedures, cohort data were linked to the hospital databases used in the geographical areas covered by the participating registries. Reference populations were all liveborn children without CAs from the same population covered by the registry, in the same birth years. The registries in Tuscany and Northern Netherlands used a random sample of their population (10% and 20%, respectively) matched by date of birth and sex. No reference children were available for the three English registries. For the registries of Paris, Norway and Malta, data on hospitalization and surgical procedures were not available for this study. Rates of hospitalisation, excluding hospital admissions associated with birth only, for children with NS were compared with those in the reference population and also to children born with any major CA previously estimated by the EUROlinkCAT project.32

Data on prescribed/dispensed medications was available by linking cohorts to local electronic prescription databases from the year 2000. Data on selected medications (antibiotics, asthma, cardiac, anti-seizure and insulin) were included in the study. Prescription data were available for 5 registries (Finland, Emilia Romagna, Tuscany, Valencian Region and Wales). Data from Wales included prescriptions for medications issued by a General Practitioner (GP), whereas data from the other four regions included medications prescribed by clinicians/GP and dispensed by a pharmacy. All registries used Anatomical Therapeutical Chemical (ATC) coding except for Wales which used Read coding which was converted to ATC coding using a look-up table in SAIL (Secure Anonymised Information Linkage Databank). No registry had information on hospital inpatient prescribing.

Statistical analysis

Each registry used a common procedure for data collection, standardisation, quality control and statistical analyses, as defined in the EUROlinkCAT project.23 Data from each registry were analysed locally by the registry using common Stata syntax scripts. Aggregated data and analytic results were uploaded to a secure Central Results Repository based at Ulster University, UK and then transferred to the research team using a secure web platform.

Kaplan-Meier survival analyses were performed by each registry to account for censoring i.e. children who were lost to follow-up due to death or emigration from the study area or who had not reached their 10th birthday by 31st December 2015. The Kaplan-Meier survival estimates with 95% confidence intervals (95%CI) from each registry were then combined in a random-effects meta-analysis using a modified method by Combescure et al.33 to obtain pooled estimates. Survival estimates were calculated at the following ages: 1 week, 4 weeks and 1, 5 and 10 years.

Similarly, each registry used a Kaplan-Meier analysis to estimate the following indicators of hospitalisation: percentage of children (i) hospitalised; (ii) hospitalised with a hospital stay longer than 10 days for term-born children; (iii) undergoing surgery. The overall estimates by age group were calculated using a random effects meta-analysis.

The median Length Of Stay in hospital (LOS) per year, defined as the number of days spent in hospital between the date of admission and the date of discharge, was calculated within each registry and random effects meta-analyses were performed using the “metamedian” package in R (version 4.0.3) for the following age groups: <1 year, 1-4 years, 5-9 years. The same method was used to obtain pooled estimates of the median age at first surgery and the median number of surgical procedures for the following age groups: <1 year, 1-4 years, and 0-4 years, as not all registries had data on children up to the child’s 10th birthday.

Finally, the proportion of children receiving a prescription per year and the median number of prescriptions per year were calculated by age groups.

**RESULTS**

Data on a total of 175 children born with NS between 1995 and 2014 came from 11 EUROCAT registries listed in Table 1.

**Survival**

A total of 15 deaths were observed in the cohort during the study period (Table 2). Twelve deaths occurred in the first year of life with an infant mortality rate of 5.4% (95% CI 1.5%-10.1%) and an additional 3 deaths were observed up to age 5. Survival estimates ranged from 98.0% (95% CI 95.9%-100.0%) at 1 week to 92.4% (95% CI 87.0%-98.1%) at 10 years.

**Hospitalisations**

Overall, 87.9% (95% CI 75.3%-94.3%) had at least one hospital stay in the first year of life (Table 3), which decreased to 65.1% per year (95% CI 45.2%-79.3%) at age 5-9 years. The percentage of children with NS who were hospitalised was three times higher than the reference population at all ages. After the first year of life, the proportion is also higher than that of children with any major CA (Figure 1). In the first year of life, 31.4% (95% CI 17.2%-46.8%) of term-born children had at least one single hospital stay longer than 10 days. The proportion decreased after the first year.

The median LOS was 15.3 days (95% CI 9.3-21.2) in children with less than 1 year of age and dramatically decreased after the first year of age.

**Surgery and other procedures**

Overall, 65.2% (95% CI 41.0%-81.5%) of children underwent surgery in the first 5 years of life and 34.8% in the first year (Table 4). The median number of surgical procedures in the first 5 years was 2 (95% CI 1.3-2.7). The median age at first surgery was 29.0 weeks (95% CI 13.6-84.4).

**Prescriptions**

About 50% of children had at least one anti-asthmatic prescription during the first 10 years of life. The percentage was quite stable across the age group (Table 5). Cardiac medications were prescribed to about 22% of children during the study period. The percentage of children with cardiac medication in the first year was 10.7% and it was 2.9% yearly in children aged 5-9. Antibiotics were prescribed to 53.6% of the children in the first year and the proportion increased with increasing age. After the first year of age, in fact, 62.4% of children had on average two prescriptions every year until 4 years of age. After four years of age, the percentage of prescriptions decreased. In the investigated cohort, only one child had an anti-seizure medication.

**DISCUSSION**

In our study, we estimated a survival rate of 92.4% at 10 years which is somewhat lower than the estimate calculated in a study on patients affected by RASopathies by Calcagni et al.17 This can be explained by the fact that our cohort consisted of children with NS diagnosed in infancy according to EUROCAT guidelines, which represented more severe cases of NS. The survival in NS is dependent on the severity of the phenotype, particularly the severity of the heart defect. This study shows that the prognosis significantly improves after one year of age.

The percentage of hospitalized children with NS was high in all age groups, especially in the early years of life. Factors that may contribute to longer hospital stays in the first years of life for patients with NS include feeding difficulties, respiratory tract infections, and the presence of congenital heart defects,.16,34-36 Cardiac abnormalities are present in most patients with NS and may require cardiac interventions, leading to extended hospital stays. Feeding problems and failure to thrive in the first year of life are very common in infants with NS. Another EUROlinkCAT study estimated that about 8% of NS patients need gastrostomy for tube feeding indicating that feeding problems may be a major cause of long hospital stays in infancy. 37

A study found that paediatric cardiac surgery admissions tended to be 4.5 days longer and cost $54,296 more in total charges for patients with NS compared to those without the syndrome. Inpatient mortality was also increased.22 Additionally, patients with NS are more likely to experience complications such as chylothorax, or perioperative bleeding from coagulation defects which can further prolong hospitalization. The increased complexity of surgical procedures and the need for concomitant procedures and re-interventions, also cause longer hospital stays.22,38-40 This helps to explain our findings that the percentage of hospitalised children with NS was higher compared to other children with or without CA in all age groups. Assessment of the total hospital care needs is important for a proper and efficient planning of healthcare services.

Individuals with NS may require surgery to address specific medical conditions or complications associated with the syndrome. Different types of surgery that may be performed in individuals with NS include cardiac surgery, airway surgery, orthognathic surgery, orchidopexy and neurosurgical interventions. Cardiac surgery may be necessary to repair or replace heart valves or correct other structural abnormalities16,38 Some individuals with NS may have airway abnormalities, such as lymphangiomatosis of the pleura, lungs, and chest wall, which can lead to complications like chylothorax. Airway surgery may be performed to address these issues.41 Corrective jaw surgery may be performed in individuals with NS who have craniofacial abnormalities. This surgery can help improve facial symmetry and correct malocclusion.42 In rare cases, individuals with NS may require neurosurgical interventions for conditions such as Chiari malformation, syringomyelia, craniosynostosis or brain tumours.43 The results of our study show surgical procedures are often performed and that two-thirds of children with NS have undergone more than one surgical procedure in the first 5 years of life. The expected frequency of surgery is an important information for healthcare providers, clinicians and parents to children with NS.

Regarding the results of the selected categories of medications investigated in the EUROlinkCAT project,23 we found that the prevalence of prescription is quite high for antibiotics in early infancy. In particular, the use of antibiotics in children with NS increased after the first year of life, whereas the rates of prescription of antibiotics in the general population estimated in some European countries decreased after the first year.44,45 This may indicate more frequent infectious diseases in children with NS at that age.

Due to the chronic nature and severity of congenital heart defects, the median number of prescriptions per year was the highest for cardiac medications. The decreasing prevalence of prescriptions of cardiac medications is expected as most children with NS have less severe congenital heart defects, such as ventricular septal defects or pulmonary valve stenosis, which rarely need any medical treatment after the first year.

**Strengths**

The main strength of this study was the population-based design which included all children diagnosed in the first year of life with NS in the areas covered by the participating EUROCAT registries and not only children referred to tertiary care centres. The population-based setting allows the calculation of more representative estimates of health outcomes. Additionally, we used data collected and validated by EUROCAT registries which use standardised definitions and coding of major CAs. Pooling data from 11 registries in 8 European countries allows powerful investigation of a rare syndrome like NS. Finally, health outcomes were estimated using an innovative methodology based on standardized procedures of analysis and the use of a meta-analytic approach.23,46

**Limitations**

A limitation of the study was that failing to link to the records in the healthcare databases could have produced bias in the estimates of some health outcomes. However, the overall successful linkage was higher than 95% and similar to that observed for the reference population (i.e. liveborn children without any congenital anomaly). Furthermore, linkage failure is most likely to occur in the first days of life before the newborns have their permanent name or Identification number as showed in a previous EUROlinkCAT study.27 As survival in the first day is high in children born with NS, this limitation is likely to have a minor impact. Another limitation of the study is that we were not able to analyse data separately by type of associated anomalies due to the very small sample size in most of the participating registries. Also, we were not able to analyse the diagnosis for the hospital stays. However, the results provide information on what parents to children with NS can expect in relation to mortality and morbidity up to 10 years of age. Finally, data on hospital inpatient prescriptions were not available for the investigated groups of medications and this might lead to a slight underestimation.

**CONCLUSIONS**

This multicentre population-based study reported on rates of survival, hospitalisations, and prescriptions of children born with NS in eleven areas of seven European countries. Survival at 1 and 10 years was 95% and 92%, respectively. In the first year of life one-third of children had a long hospital stay. After the first year, hospitalisation was still high, but the median length of stay dramatically decreased. Two-thirds of children had a surgical procedure in the first five years of life with a medium of two interventions. The results of the study provide information to evaluate the health burden for children diagnosed with NS that is important for parental counselling and the planning of healthcare services. Efforts must be made to support families throughout the childhood of children born with NS.

**Declarations**

**Ethics approval and consent to participate**

All EUROCAT registries contributing data to the EUROlinkCAT project obtained ethical, governance and other permissions for the data linkage according to their national legislations and arrangements. University of Ulster obtained Ethics permission for the Central Results Repository on 15 September 2017 (Institute of Nursing and Health Research Ethics Filter Committee, number FCNUR-17-000).

**Consent for publication**

The work described has not been submitted elsewhere for publication, in whole or in part, and all the authors listed have approved the manuscript for publication.

**Availability of data and materials**

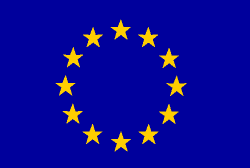
The data that support the findings of this study are available from the participating registries of congenital anomalies, but restrictions apply to the availability of these data, which were used under license for the current study, and so are not publicly available. Data are however available from the authors for scientifically valid requests and with permission of the participating registries of congenital anomalies.

**Competing interest**

The authors have no conflicts of interest relevant to this article to disclose.

**Funding/Support**

This project has received funding from the European Union’s Horizon 2020 research and innovation programme under grant agreement No. 733001 (Jan 2017 – Dec 2021) <https://ec.europa.eu/programmes/horizon2020/en>).



The funder had no role in the design and conduct of the study; collection, management, analysis, and interpretation of the data; preparation, or review; and decision to submit the manuscript for publication. The views presented here are those of the authors only, and the European Commission is not responsible for any use that may be made of the information presented here.

**Authors contributions**

MS conceptualized and designed the study, developed statistical analysis plan, performed the analysis, interpreted the results, drafted the initial manuscript, and reviewed and revised the manuscript. IB conceptualized and designed the study, interpreted the results, drafted the initial manuscript, and reviewed and revised the manuscript. AC conceptualized and designed the study, developed statistical analysis plan, contributed to the data analysis and interpretation of the results, and critically reviewed the manuscript for important intellectual content. JT contributed to development of study methods, including data standardization and data linkage, development of statistical analysis plan, writing analysis programs, data analysis and interpretation of the results, and critically reviewed the manuscript for important intellectual content. EG contributed to obtaining funding, development of study methods, including data standardization and data linkage, to interpretation of the results, and critically reviewed the manuscript for important intellectual content. ML contributed to obtaining funding, development of study methods, was responsible for data standardization and management of data linkage by the participating data providers, contributed to data analysis and interpretation of the results, and critically reviewed the manuscript for important intellectual content. LO and MVA contributed to design the study, interpreted the results and critically reviewed the manuscript for important intellectual content. EB, CC, MGa, MGi, KK, NL, DT,and DW were responsible for data linkage and standardization for their registries’ data and running centrally written syntax scripts for local analyses, and critically reviewed the manuscript for important intellectual content. JM conceptualized and designed the study, obtained funding, developed study methods, including data standardization and linkage, supervised writing analysis programs, performed statistical analysis, supervised the work, and reviewed and revised the manuscript.

All authors approved the final manuscript as submitted and agree to be accountable for major aspects of the work.

**Aknowledgements:** Not applicable

**Abbreviations:** NS**:** Noonan syndrome; HCM: hypertrophic cardiomyopathy; CA: congenital anomalies; ICD-10 BPA: International Statistical Classification of Diseases, Tenth Revision, with British Paediatric Association 1- digit extension; ICD-9 BPA: International Statistical Classification of Diseases, Ninth Revision, with British Paediatric Association 1- digit extension; LOS: length of stay in hospital ATC: Anatomical Therapeutical Chemical; SAIL: Secure Anonymised Information Linkage

**REFERENCES**

1. Tiemens DK, Kleimeier L, Leenders E, et al. The most important problems and needs of rasopathy patients with a noonan syndrome spectrum disorder. Orphanet J Rare Dis. 2023 Jul 21;18(1):198. doi: 10.1186/s13023-023-02818-y.
2. Hebron KE, Hernandez ER, Yohe ME. The RASopathies: from pathogenetics to therapeutics. Dis Model Mech. 2022 Feb 1;15(2):dmm049107. doi: 10.1242/dmm.049107. Epub 2022 Feb 18.
3. Zenker M. Clinical overview on RASopathies. Am J Med Genet C Semin Med Genet. 2022 Dec;190(4):414-424. doi: 10.1002/ajmg.c.32015. Epub 2022 Nov 25.
4. Shaw AC, Kalidas K, Crosby AH, et al. The natural history of Noonan syndrome: a long-term follow-up study. Arch Dis Child. 2007 Feb;92(2):128-32. doi: 10.1136/adc.2006.104547.
5. Van der Burgt I. Noonan syndrome. Orphanet J Rare Dis. 2007 Jan 14;2:4. doi: 10.1186/1750-1172-2-4
6. Romano AA, Allanson JE, Dahlgren J, et al. Noonan syndrome: clinical features, diagnosis, and management guidelines. Pediatrics. 2010 Oct;126(4):746-59. doi: 10.1542/peds.2009-3207.
7. Roberts AE, Allanson JE, Tartaglia M, Gelb BD. Noonan syndrome. Lancet. 2013 Jan 26;381(9863):333-42. doi: 10.1016/S0140-6736(12)61023-X.
8. Capri Y, Flex E, Krumbach OHF, et al. Activating Mutations of RRAS2 Are a Rare Cause of Noonan Syndrome. Am J Hum Genet. 2019 Jun 6;104(6):1223-1232. doi: 10.1016/j.ajhg.2019.04.013.
9. Roberts AE. Noonan Syndrome. 2001 Nov 15 [updated 2022 Feb 17]. In: Adam MP, Feldman J, Mirzaa GM, et al. editors. GeneReviews® [Internet]. Seattle (WA): University of Washington, Seattle; 1993–2023.
10. Zenker M, Edouard T, Blair JC, Cappa M. Noonan syndrome: improving recognition and diagnosis. Arch Dis Child. 2022 Dec;107(12):1073-1078. doi: 10.1136/archdischild-2021-322858
11. Allen MJ, Sharma S. Noonan Syndrome. 2023 Jan 9. In: StatPearls [Internet]. Treasure Island (FL): StatPearls Publishing; 2023 Jan–.
12. Briggs BJ, Dickerman JD. Bleeding disorders in Noonan syndrome. Pediatr Blood Cancer. 2012 Feb;58(2):167-72. doi: 10.1002/pbc.23358.
13. Nugent DJ, Romano AA, Sabharwal S, Cooper DL. Evaluation of bleeding disorders in patients with Noonan syndrome: a systematic review. J Blood Med. 2018 Oct 23;9:185-192. doi: 10.2147/JBM.S164474.
14. Lioncino M, Monda E, Verrillo F, Moscarella E, Calcagni G, Drago F, Marino B, Digilio MC, Putotto C, Calabrò P, Russo MG, Roberts AE, Gelb BD, Tartaglia M, Limongelli G. Hypertrophic Cardiomyopathy in RASopathies: Diagnosis, Clinical Characteristics, Prognostic Implications, and Management. Heart Fail Clin. 2022 Jan;18(1):19-29. doi: 10.1016/j.hfc.2021.07.004.
15. Edouard T, Zenker M, Östman-Smith I, Ortega Castelló E, Wolf CM, Burkitt-Wright E, Verloes A, García-Miñaúr S, Tartaglia M, Shaikh G, Lebl J. Management of growth failure and other endocrine aspects in patients with Noonan syndrome across Europe: A sub-analysis of a European clinical practice survey. Eur J Med Genet. 2022 Jan;65(1):104404. doi: 10.1016/j.ejmg.2021.104404
16. Linglart L, Gelb BD. Congenital heart defects in Noonan syndrome: Diagnosis, management, and treatment. Am J Med Genet C Semin Med Genet. 2020 Mar;184(1):73-80. doi: 10.1002/ajmg.c.31765. Epub 2020 Feb 5. PMID: 32022400; PMCID: PMC7682536.
17. Calcagni G, Limongelli G, D'Ambrosio A, et al. Cardiac defects, morbidity and mortality in patients affected by RASopathies. CARNET study results. Int J Cardiol. 2017 Oct 15;245:92-98. doi: 10.1016/j.ijcard.2017.07.068.
18. Meier AB, Raj Murthi S, Rawat H, et al. Cell cycle defects underlie childhood-onset cardiomyopathy associated with Noonan syndrome. iScience. 2021 Dec 9;25(1):103596. doi: 10.1016/j.isci.2021.103596.
19. Hickey EJ, Mehta R, Elmi M, et al. Survival implications: hypertrophic cardiomyopathy in Noonan syndrome. Congenit Heart Dis. 2011 Jan-Feb;6(1):41-7. doi: 10.1111/j.1747-0803.2010.00465.x
20. Wilkinson JD, Lowe AM, Salbert BA, et al. Outcomes in children with Noonan syndrome and hypertrophic cardiomyopathy: a study from the Pediatric Cardiomyopathy Registry. Am Heart J. 2012 Sep;164(3):442-8. doi: 10.1016/j.ahj.2012.04.018.
21. Kaltenecker E, Schleihauf J, Meierhofer C, Shehu N, Mkrtchyan N, Hager A, Kühn A, Cleuziou J, Klingel K, Seidel H, Zenker M, Ewert P, Hessling G, Wolf CM. Long-term outcomes of childhood onset Noonan compared to sarcomere hypertrophic cardiomyopathy. Cardiovasc Diagn Ther. 2019 Oct;9(Suppl 2):S299-S309. doi: 10.21037/cdt.2019.05.01.
22. Kriz C, Flores S, Villarreal EG, et al. Impact of Noonan Syndrome on admissions for pediatric cardiac surgery. Minerva Pediatr (Torino). 2022 Aug;74(4):461-467. doi: 10.23736/S2724-5276.19.05461-6. Epub 2019 Jun 28.
23. Morris JK, Garne E, Loane M, et al. EUROlinkCAT protocol for a European population-based data linkage study investigating the survival, morbidity and education of children with congenital anomalies. 2021 Jun 28;11(6):e047859. doi: 10.1136/bmjopen-2020-047859.
24. Boyd PA, Haeusler M, Barisic I, et al. Paper 1: The EUROCAT network-organization and processes. Birth Defects Res A Clin Mol Teratol. 2011 Mar;91 Suppl 1:S2-15. doi: 10.1002/bdra.20780.
25. Kinsner-Ovaskainen A, Lanzoni M, Garne E, et al., A sustainable solution for the activities of the European network for surveillance of congenital anomalies: EUROCAT as part of the EU Platform on Rare Diseases Registration. Europ J Med Genetics, 2018. 61(9): p. 513-517.
26. EUROCAT Guide 1.5: https://eu-rd-platform.jrc.ec.europa.eu/eurocat/data-collection/guidelines-for-data-registration\_en#inline-nav-2
27. Loane M, Given JE, Tan J et al. Linking a European cohort of children born with congenital anomalies to vital statistics and mortality records: A EUROlinkCAT study. PLOS ONE. 2021;16(8):e0256535. doi: 10.1371/journal.pone.0256535
28. Loane M, Given JE, Tan J, et al. Creating a population-based cohort of children born with and without congenital anomalies using birth data matched to hospital discharge databases in 11 European regions: Assessment of linkage success and data quality. PLoS One. 2023 Aug 30;18(8):e0290711. doi: 10.1371/journal.pone.0290711.
29. Glinianaia SV, Rankin J, Pierini A, et al. Ten-Year Survival of Children With Congenital Anomalies: A European Cohort Study. Pediatrics. 2022 Feb 11:e2021053793. doi: 10.1542/peds.2021-053793
30. Santoro M, Coi A, Pierini A, et al. Temporal and geographical variations in survival of children born with congenital anomalies in Europe: A multi-registry cohort study. Paediatr Perinat Epidemiol. 2022 Nov;36(6):792-803. doi: 10.1111/ppe.12884
31. Coi A, Santoro M, Pierini A, et al. Survival of children with rare structural congenital anomalies: a multi-registry cohort study. Orphanet J Rare Dis. 2022 Mar 29;17(1):142. doi: 10.1186/s13023-022-02292-y

Urhoj SK, Tan J, Morris JK, Given J, et al. Hospital length of stay among children with and without congenital anomalies across 11 European regions-A population-based data linkage study. PLoS One. 2022 Jul 22;17(7):e0269874. doi: 10.1371/journal.pone.0269874.

Combescure C, Foucher Y, Jackson D. Meta-analysis of single-arm survival studies: a distribution-free approach for estimating summary survival curves with random effects. Stat Med. 2014;33(15):2521-2537.

Croonen EA, Draaisma JMT, van der Burgt I, et al. First-year growth in children with Noonan syndrome: Associated with feeding problems? Am J Med Genet A. 2018 Apr;176(4):951-958. doi: 10.1002/ajmg.a.38649.

Sleutjes J, Kleimeier L, Leenders E, et al. Lymphatic Abnormalities in Noonan Syndrome Spectrum Disorders: A Systematic Review. Mol Syndromol. 2022 Feb;13(1):1-11. doi: 10.1159/000517605

Pieper CC, Wagenpfeil J, Henkel A, et al. MR lymphangiography of lymphatic abnormalities in children and adults with Noonan syndrome. Sci Rep. 2022 Jul 1;12(1):11164. doi: 10.1038/s41598-022-13806-w.

Garne E, Tan J, Loane M, et al. Gastrostomy and congenital anomalies: a European population-based study. BMJ Paediatrics Open 2022;6:e001526. doi:10.1136/bmjpo-2022-001526

Hemmati P, Dearani JA, Daly RC, et al. Early Outcomes of Cardiac Surgery in Patients with Noonan Syndrome. Semin Thorac Cardiovasc Surg. 2019 Autumn;31(3):507-513. doi: 10.1053/j.semtcvs.2018.12.004

1. Morice A, Harroche A, Cairet P, Khonsari RH. Preoperative Detailed Coagulation Tests Are Required in Patients With Noonan Syndrome. J Oral Maxillofac Surg. 2018 Jul;76(7):1553-1558. doi: 10.1016/j.joms.2017.12.012.
2. Holzmann J, Tibby SM, Rosenthal E, et al. Results of balloon pulmonary valvoplasty in children with Noonan's syndrome. Cardiol Young. 2018 May;28(5):647-652. doi: 10.1017/S1047951117002827.
3. Yellon RF. Complications following airway surgery in Noonan syndrome. Arch Otolaryngol Head Neck Surg. 1997 Dec;123(12):1341-3. doi: 10.1001/archotol.1997.01900120091015.
4. Lutz JC, Nicot R, Schlund M, et al. Dental and maxillofacial features of Noonan Syndrome: Case series of ten patients. J Craniomaxillofac Surg. 2020 Mar;48(3):242-250. doi: 10.1016/j.jcms.2020.01.011.
5. Saragosti E, Fattal-Valevski A, Levin D, et al. Neurosurgical aspects of Noonan syndrome. Childs Nerv Syst. 2023 Apr;39(4):849-856. doi: 10.1007/s00381-023-05888-2.
6. Pottegård A, Broe A, Aabenhus R, Bjerrum L, Hallas J, Damkier P. Use of antibiotics in children: a Danish nationwide drug utilization study. Pediatr Infect Dis J. 2015 Feb;34(2):e16-22
7. Dekker ARJ, Verheij TJM, van der Velden AW. Antibiotic management of children with infectious diseases in Dutch Primary Care. Fam Pract. 2017 Apr 1;34(2):169-174. doi: 10.1093/fampra/cmw125.
8. Gilboa SM, Tepper NK, Reefhuis J. Multijurisdictional Analyses of Birth Defects: Considering the Common Data Model Approach. Pediatrics. 2022;149(3):e202105528

**TABLES AND FIGURES**

**Table 1.** Contributing European Surveillance of Congenital Anomalies (EUROCAT) registries, included birth years

|  |  |
| --- | --- |
| **Participating registries** | **Included birth years** |
| Finlande | 1995-2014b |
| France: Parisa,d | 1995-2014 |
| Italy: Emilia Romagna | 2008-2014 |
| Italy: Tuscany | 2005-2014 |
| Maltaa,d,e | 1995-2014 |
| Norwaya,d,e | 1999-2014 |
| Spain: Valencian Region | 2007-2014c |
| UK: East Midlands and South Yorkshired | 2003-2012 |
| UK: Thames Valleyd | 2005-2013 |
| UK: Walese | 1998-2014 |
| UK: Wessexd | 2004-2014 |

a data on hospitalisation and surgical procedures not available

b study period for hospitalisation and surgical procedures 1997-2014

c study period for hospitalisation and surgical procedures 2010-2014

d data on prescriptions not available

f whole nation covered

**Table 2**. Pooled survival estimates at selected age groups up to 10 years of age for children born with Noonan syndrome (n=175), 1995-2014

|  |  |  |  |
| --- | --- | --- | --- |
| **Age** | **N. of deaths** | **Survival %** | **95% CI** |
| 1 week | 4 | 98.0 | 95.9-100.0 |
| 4 weeks | 6 | 96.7 | 93.9-99.6 |
| 1 year | 12 | 94.6 | 90.9-98.5 |
| 5 years | 15 | 92.6 | 87.4-98.1 |
| 10 years | 15 | 92.4 | 87.0-98.1 |

**Table 3**. Percentage hospitalised, percentage hospitalised with a long stay (≥10 days) and median length of stay per year of children with Noonan syndrome (n=145), by age, 1995-2014

|  |  |  |  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- | --- | --- | --- |
| **Age (year)** | **Children with any hospitalisation** | | | **Children with a length of stay over 10 daysa** | | | **Median length of stay** | |
| **n** | **%** | **95% CI** | **n** | **%** | **95% CI** | **days** | **95% CI** |
| **<1** | 126 | 87.9 | 75.3-94.3 | 35 | 31.4 | 17.2-46.8 | 15.3 | 9.3-21.2 |
| **1-4** | 104 | 78.5 | 64.1-87.7 | 17 | 15.8 | 3.2-37.0 | 1.3 | 0.3-2.2 |
| **5-9** | 52 | 65.1 | 45.2-79.3 | 3 | - | - | 0.5 | 0.0-1.0 |

**a** Only children born ≥37 weeks of gestation were included.

**Figure 1**. Percentage hospitalised (with 95% CIs) children with Noonan syndrome, children with any congenital anomaly (‘Any CA’) and children without a congenital anomaly (‘Reference’), by age, 1995-2014

**Table 4.** Proportion of children with Noonan syndrome (n=145) undergoing surgery and median number of surgeries by age, 1995-2014

|  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- |
| **Age (year)** | **N children undergoing surgery** | **% children undergoing surgery** | **95% CI** | **Median number of surgeries** | **95% CI** |
| **<1** | 50 | 34.8 | 21.1-49.0 | 1.3 | 0.7-1.8 |
| **1-4** | 60 | 50.8 | 22.3-73.7 | 2.0 | 1.7-2.3 |
| **0-4** | 80 | 65.2 | 41.0-81.5 | 2.0 | 1.3-2.7 |

**Table 5**. Percentage of children with a prescription and median number of prescriptions by age and type of prescription, 2000-2014

|  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- |
| **Medication** | **Age (year)** | **Percentage of children with a prescription in a year** | **95% CI** | **Median number of prescriptions each year** | **IQR** |
| **Any asthma medication**  (ATC code R03) | **<1** | 17.9 | 11.3-26.2 | 2 | 2-2 |
| **1-4** | 24.7 | 20.5-29.4 | 2 | 1-4 |
| **5-9** | 18.2 | 13.8-23.2 | 3.5 | 2-5 |
| **Any cardiac medication**  (ATC code C01-C03, C07-C09, excluding C01BA51, C01BA71, C01CA24) | **<1** | 10.7 | 5.7-18.0 | 8.5 | 5.5-9.5 |
| **1-4** | 8.4 | 5.8-11.7 | 6 | 4-10 |
| **5-9** | 2.9 | 1.2-5.5 | 4.5 | 4-5 |
| **Antibacterials for systemic use**  (ATC code J01-J05) | **<1** | 53.6 | 43.9-63.0 | 2 | 1-4 |
| **1-4** | 62.4 | 57.3-67.3 | 2 | 1-4 |
| **5-9** | 38.4 | 32.7-44.4 | 1.5 | 1-3 |

IQR= Interquartile range