Resolving the Genomic Complexity of Pediatric Acute Lymphoblastic Leukemia

YANARA MARINCEVIC-ZUNIGA
Acute lymphoblastic leukemia (ALL) is the most common pediatric cancer in the Nordic countries. Structural chromosomal rearrangements are a hallmark of ALL and represent key markers for diagnosis, risk stratification and prognosis. Nevertheless, a substantial proportion of ALL cases (~25%) lack known risk-stratifying markers and are commonly referred to as the B-other subgroup. Improved delineation of structural alterations within this subgroup could provide additional information for diagnosis, prognosis and treatment decisions. Therefore, the aim of this thesis was to decipher the genetic alterations in pediatric ALL, focusing on patients in the B-other subgroup that lack known risk-stratifying markers, and to gain further understanding of the prognostic relevance of aberrant chromosomal changes in ALL.

This thesis comprises four studies. In study I we identified a novel and recurrent fusion gene (PAX5-ESRRB) in four B-other patients using a combination of RNA-sequencing and copy number analysis. These patients displayed a distinct gene expression and DNA-methylation pattern that differed from other subtypes of ALL. In study II we further explored the fusion gene landscape in ALL by applying RNA-sequencing to 134 patient samples assigned to different subtypes, including the B-other subgroup. We detected several novel and recurrent fusion gene families in approximately 80% of the B-other patients of which several were associated with distinct DNA methylation and gene expression profiles. Following on from study II, in study III we utilized subtype-specific DNA methylation patterns to design DNA methylation-based classifiers to screen for subtype membership in ~1100 ALL samples including a large group of B-other samples (25%). Re-classification of B-other samples into a new subtype using DNA methylation as the sole marker for subtype classification was validated by RNA-sequencing, which identified previously unknown fusion genes. In study IV, “linked-read” whole genome sequencing was applied to 13 ALL samples for in-depth analysis of chromosomal rearrangements. We detected all known pathogenic variants with this technique and also identified previously unknown structural aberrations at a resolution beyond that obtained by traditional karyotyping.

Together, these studies provide novel insights into the structural variation present in ALL and their potential clinical relevance, which may contribute to improved treatment stratification and risk-evaluation of children diagnosed with ALL in the future.

Keywords: Acute lymphoblastic leukemia (ALL), 450k array, DNA methylation, RNA-sequencing, linked-read WGS, B-other, fusion gene, chromosomal translocation

Yanara Marincevic-Zuniga, Department of Medical Sciences, Molecular Medicine, Akademiska sjukhuset, Uppsala University, SE-75185 Uppsala, Sweden.

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Dedicado a mi familia
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List of Papers

This thesis is based on the following papers, which are referred to in the text by their Roman numerals.

I  

II  

III  

IV  

*Equally contributing authors.

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## Abbreviations

<table>
<thead>
<tr>
<th>Abbreviation</th>
<th>Definition</th>
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<tbody>
<tr>
<td>ALL</td>
<td>Acute lymphoblastic leukemia</td>
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<tr>
<td>BCP-ALL</td>
<td>B-cell precursor acute lymphoblastic leukemia</td>
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<tr>
<td>chr</td>
<td>Chromosome</td>
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<td>CIMP</td>
<td>CpG island methylator phenotype</td>
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<td>CNA</td>
<td>Copy number alteration</td>
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<td>CpG</td>
<td>CG dinucleotide</td>
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<td>dic(9;20)</td>
<td>Dicentric rearrangement between chromosomes 9 and 20</td>
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<td>DMC</td>
<td>Differentially methylated CpG</td>
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<td>DNMT</td>
<td>DNA methyltransferase</td>
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<td>FISH</td>
<td>Fluorescence in situ hybridization</td>
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<tr>
<td>FPKM</td>
<td>Fragments per kilobase per million mapped reads</td>
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<tr>
<td>HeH</td>
<td>High hyperdiploid</td>
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<tr>
<td>HMW DNA</td>
<td>High-molecular weight DNA</td>
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<tr>
<td>iAMP21</td>
<td>Intrachromosomal amplification of chromosome 21</td>
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<td>Ig</td>
<td>Immunoglobulin</td>
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<td>MRD</td>
<td>Minimal residual disease</td>
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<td>NGS</td>
<td>Next-generation sequencing</td>
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<td>NOPHO</td>
<td>Nordic Society of Pediatric Hematology and Oncology</td>
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<tr>
<td>PCA</td>
<td>Principal component analysis</td>
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<td>RACE</td>
<td>Rapid amplification of cDNA ends</td>
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<td>RIN</td>
<td>RNA integrity number</td>
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<tr>
<td>RT-PCR</td>
<td>Reverse transcription polymerase chain reaction</td>
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<td>SNP</td>
<td>Single nucleotide polymorphism</td>
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<td>SNV</td>
<td>Single nucleotide variant</td>
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<td>T-ALL</td>
<td>T-cell lineage acute lymphoblastic leukemia</td>
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<td>TAPS</td>
<td>Tumor Aberration Prediction Suite</td>
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<td>TCR</td>
<td>T-cell receptor</td>
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<td>WGS</td>
<td>Whole genome sequencing</td>
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Introduction

The hematopoietic stem cell is the founder unit to all the cells in the blood system. These cells undergo a series of hierarchical differentiation steps during hematopoiesis, giving rise to a wide plethora of mature blood cells. Leukemia is a heterogeneous malignant disorder that arises upon the acquisition of genetic alterations in immature blood cells during different stages of the hematopoietic hierarchy. These changes perturb multiple cellular pathways, ultimately leading to abnormal cellular expansion of leukemic cells. Genomic profiling to uncover the pathogenic alterations underlying leukemic cell development is essential in order to understand which of the molecular features that are of clinical relevance. This thesis focuses on using advanced technologies for identifying factors that may have a functional role in the pathogenesis of a specific sub-class of lymphoid leukemia.

Acute lymphoblastic leukemia

Acute lymphoblastic leukemia (ALL) is a disease of the white blood cells that arises from the early B-cell precursor (BCP-ALL) or T-cell (T-ALL) lineages in the bone marrow. The malignant transformation triggers a rapid clonal expansion of the leukemic cells, which out-competes the normal functioning cells, and ultimately disseminate into the peripheral blood system. The term “acute” in ALL reflects the quick progression of this transformation, and if not treated immediately, the outcome is dismal. While relatively uncommon in adults, ALL is the most common cancer amongst children with a peak age of incidence between two and five years of age [1]. Each year, there are approximately 200 children diagnosed with ALL in the Nordic countries.

Although ALL is a malignant disease, the modern advancements in combination therapy and improved diagnostics have been a true success story. Today over 90% of ALL patients reach complete clinical remission after induction therapy [2, 3], which is in contrast to the approximately 20-40% overall survival rates reported during the 1960-1970s [4-6]. However, despite the improved outcome for ALL patients, 15-20% of patients still suffer from relapse or do not respond to treatment, and are the main reasons for ALL-related mortality in children [7]. Relapse in ALL is typically associated with a much poorer outcome, however, recent studies have shown that the 5-
year overall survival rate for relapsed ALL cases has improved over the last two decades reaching over 50% [7]. Together, these marked improvements can be attributed to several landmarks within treatment advancement, such as the discovery of targeted therapies using tyrosine kinase inhibitors, but also several crucial biological findings [4].

In this introduction, I will highlight some of the key genetic findings throughout the modern history of ALL that have increased our knowledge of the underlying biology and further aided the advancement of successful treatment in ALL.

**Cytogenetic subtypes of ALL**

ALL is genetically a heterogeneous disease that is broadly classified into two immunophenotypic groups based on the cell of origin. ALL originating from the B-cell lineage is the most frequent immunophenotype accounting for almost 85% of the cases, while only 15% of the ALL cases arise from the T-cell lineage. At the molecular level, ALL is characterized by a variety of distinct genetic abnormalities. BCP-ALL is further subdivided into several cytogenetic subtypes that are observed in varying frequencies (Figure 1). In general, BCP-ALL can be divided into four different subgroupings: 1) cases with abnormal chromosomal numbers (aneuploidies); 2) cases with translocations giving rise to recurrent and expressed fusion genes; 3) additional recognized chromosomal abnormalities; 4) genetically unclassified cases with or those without any detectable cytogenetic aberration.

![Figure 1. Distribution of risk-stratifying subtypes and B-other in BCP-ALL patients diagnosed in the Nordic countries between 1996 and 2012.](image)
BCP-ALL

Aneuploidies
Numerical chromosomal copy number alterations (aneuploidies) are frequently detected in ALL. In fact, the most common subtype is characterized by the presence of 51-67 chromosomes per leukemic blast cell and is termed high hyperdiploid (HeH). This subtype accounts for approximately 25-30% of the diagnosed pediatric BCP-ALLs. The acquisition of extra chromosomes is a non-random event and may involve any chromosome, but typical patterns of trisomies are seen for chromosomes X, 4, 6, 10, 14, 17, 18, and 21 [8]. Gains of chromosome 21, manifested by one to five extra copies, are frequent in HeH and detected in over 90% of the cases [9]. In contrast to HeH, a much rarer subtype present in approximately 1% of ALL, is defined by <45 chromosomes and termed hyperdiploid.

Translocations
Chromosomal rearrangements giving rise to expressed fusion genes are a hallmark of ALL. The most recurrent translocations used for cytogenetic classification are t(12;21), t(1;19), t(9;22) and translocations involving KMT2A (referred to as the 11q23/MLL subtype throughout this thesis).

The second most common subtype in ALL is defined by the t(12;21) translocation giving rise to the ETV6-RUNXI fusion gene, which is detected in approximately 25% of BCP-ALL. RUNXI is an essential transcription factor for definitive hematopoiesis, supported by the lack of primary hematopoiesis during embryogenesis in RUNXI knock-out mice models [10]. ETV6 also plays an important role in later stages of hematopoiesis and acts as a transcriptional repressor [11, 12]. The current model for the disease triggering mechanism of t(12;21)ETV6-RUNXI implies a two-hit process, where a prenatally acquired fusion gene alone does not have the ability to develop overt ALL, and hence additional mutation(s) are required [13, 14]. Such a two-hit model is supported by several findings such as detection of t(12;21)ETV6-RUNXI in neonatal blood spots from children that developed ETV6-RUNXI-positive ALL several years after birth, even after a latency period up to 14 years, and a low concordance rate of ETV6-RUNXI-positive ALL in monozygotic twins (5%) [15, 16]. Interestingly, studies have shown that t(12;21)ETV6-RUNXI occurs frequently during normal fetal development and is observed 1-5% of healthy newborns, although very few of these children develop ALL, which further supports the low oncogenic potential of this fusion gene [15, 17].

Two subtypes that historically have been associated with dismal prognosis are the t(1;19)TCF3-PBX1 and the t(9;22)BCR-ABL1 subtypes. Together, these subtypes represent approximately 6-8% of the BCP-ALL cases. The well described t(9;22) translocation, also known as the Philadelphia chromosome, gives rise to the chimeric protein BCR-ABL1 that functions as a con-
stitutively active tyrosine kinase. Presence of BCR-ABL1 leads to aberrant phosphorylation of various downstream target proteins and thus to the activation of multiple pathways that promote cell survival and growth [18]. The t(1;19) translocation causes the disruption and joining of two important transcription factors TCF3 and PBX1, that are regulators of normal hematopoiesis [19, 20]. The previously poor prognosis associated with t(1;19)TCF3-PBX1 has markedly improved by contemporary treatment [21]. Similarly, the discovery of tyrosine kinase inhibitors, such as Imatinib, has led to a substantial improvement of the outcome for patients with t(9;22)BCR-ABL1 and is considered a paradigm for targeted cancer therapies [22].

Rearrangement of the KMT2A gene (MLL) is the genetic alteration that defines the 11q23/MLL subtype, which is present in approximately 5% of the ALL cases. The KMT2A gene is a promiscuous gene known to form fusion genes with over 80 gene partners, including the most common partner AFF1 (AF4), MLLT3 (AF9), MLLT1 (ENL) and MLLT10 (AF10) [23]. KMT2A encodes a histone methyltransferase that methylates lysine residues of the histone H3, and plays an essential role in regulating gene expression by chromatin remodeling [24]. It has been proposed that the KMT2A-related fusion genes are prenatally acquired and are by themselves sufficient for development of overt leukemia, which is supported by the young age of disease onset (typically <1 year) and a close to 100% concordance rate for BCP-ALL between monozygotic twins [25].

Structural abnormalities of specific chromosomes

In contrast to the aforementioned subtypes, the intrachromosomal amplification of chromosome 21 (iAMP21) and dicentric rearrangements involving chromosome 9 and 20 (dic(9;20)), represent more recently identified subtypes that are now recognized as recurrent and non-random events in ALL. iAMP21 is characterized by the amplification of a 5.1 Mb region on chromosome 21q involving the RUNX1 gene [26]. This subtype, constituting roughly 2% of the cases, was initially discovered by routine FISH analysis of the t(12;21)ETV6-RUNX1 subtype, reveling an absence of t(12;21) and instead several additional copies of RUNX1 [27]. iAMP21 is thus defined as having a total of 5 or more copies of RUNX1 per leukemic cell by FISH analysis [28]. The dic(9;20) subtype encompasses approximately 3% of the cases and is characterized by a dicentric chromosome resulting from the joining of the chr9p and chr20q with the retention of both centromeres from the respective chromosomes [29, 30]. In some cases, the breakpoint in chromosome 9 has been located within the PAX5 gene, which results in expressed fusion genes together with a handful of gene partners on chromosome 20, such as NOL4L, ASXL1, and KIF3B, however not all cases result in an expressed fusion gene [31].
Genetically unclassified BCP-ALL (B-other)

Despite the well-defined cytogenetic classification in BCP-ALL, up to 30% of the BCP-ALL cases remain unclassified, and the underlying genetic alterations are either unknown or non-recurrent. This group of ALL patients is collectively referred to as B-other. With the aid of several high-throughput techniques for genome-wide profiling, such as whole genome sequencing (WGS), transcriptome sequencing and array-based copy number analysis, several alterations have recently been uncovered in B-other and have reshaped our notion about this subgroup [32]. In general these findings comprise key alterations defined by distinct gene expression patterns, novel fusion genes and recurrent micro-deletions in genes affecting key pathways, as will be outlined later in this thesis.

T-ALL

Opposite to the well-defined cytogenetic classification of BCP-ALL, T-ALL is characterized by few recurrent chromosomal abnormalities and a more heterogeneous genetic landscape [33]. The most frequent alterations in T-ALL are activating NOTCH1 mutations and 9p deletions resulting in inactivation of the CDKN2A/B gene [34]. Together, these two types of alterations occur in more than 50% of the T-ALL cases. Chromosomal rearrangements giving rise to expressed fusion genes are also frequent in T-ALL, commonly involving the T-cell receptor gene [35]. Although the molecular basis of T-ALL has radically been redefined in recent years, most of the described genetic alterations have had a low prognostic impact [33].

Risk-associated markers and clinical management in ALL

As the multiple ALL subtypes are associated with different clinical outcome the discovery of recurrent genetic abnormalities constitute yet another landmark in the clinical management of ALL (Figure 2). Thus, the presence of genetic alterations is an important factor for risk-based stratification and directed treatments in ALL [36]. Genetic markers for “standard” risk comprise the two largest subtypes, HeH and t(12;21)ETV6-RUNXI, that are associated with favorable outcome. Three markers that confer intermediate risk are the (1;19)TCF3-PBX, dic(9;20) and the iAMP21 subtypes. Subtypes that are considered high-risk genetic markers and are associated with inferior prognosis include the less frequent subtypes t(9;22)BCR-ABL1, 11q23/MLL and hypodiploid subtypes. These subtypes are associated with higher rates of relapse, therapy resistance or death as compared to other subtypes [5].
Apart from molecular-based stratification, clinical parameters are essential for risk-associated treatment stratification of ALL [37]. An age less than one year or more than 10 years at diagnosis are considered high-risk features, which can be attributed to the age distribution amongst the ALL subtypes [5]. For instance, the 11q23/MLL subtype which confers poor prognosis is frequently observed in cases younger than 1 year whereas iAMP21 cases tend to be older, with a median age of 9. Other features such as high white blood cell count and involvement of the central nervous system are also considered to be high-risk features. Additionally, establishment of routine monitoring of minimal residual disease (MRD) to assess overall therapy response has proven to be a highly reliable prognostic indicator [38, 39]. Evaluation of MRD by flow cytometry or real-time quantitative PCR to monitor treatment response at a submicroscopic level enables detection of residual leukemic cells during or after induction therapy. Detection of MRD can be performed by screening of leukemia-specific immunophenotypes or clonal rearrangements of the T-cell receptor (TCR) or immunoglobulin (Ig) [40]. Additionally, studies have shown that monitoring of treatment response is also possible by screening of risk-associated markers, such as the ETV6-RUNXI or TCF3-PBXI fusion genes, which complements current MRD methods [41, 42]. These findings highlight the importance of establishing the underlying alteration in the leukemic cells at diagnosis, which may become fundamental markers for MRD.

Another important aspect for the clinical management of ALL has been the establishment of several international collaborative working groups, including the Nordic Society of Pediatric Hematology and Oncology group (NOPHO) across the Nordic countries that aim for unifying diagnostics and common treatment protocols and strategies [43]. As a result, intensification of treatment protocols and addition of new drugs, such as tyrosine kinase inhibitors for treatment of t(9;22)BCR-ABL1 positive ALL, have also had an impact on clinical management of ALL.
Figure 2. Kaplan-Meier curves showing the overall survival of >900 samples from the different subtypes of ALL treated according to the NOPHO protocols between 1996 and 2011.

ALL in the “omics”-era and discovery of new subtypes

The low-resolution methods such as karyotyping and FISH that are routinely used for clinical diagnostics in ALL are limited in their ability to detect novel or discreet changes of the genome, including changes at base pair resolution. Microarray-based technologies and next generation sequencing (NGS) have greatly increased our ability to identify novel submicroscopic alterations in a genome-wide context. As a consequence of such technological advancements, new molecularly distinct subgroups have been described in ALL, predominantly in the heterogeneous B-other group [32, 44, 45]. For this reason, the classification system of BCP-ALL is presently changing as several new and recurrent putative subgroups have emerged that are transforming the conventional view of subtype classification in ALL.

Genomic characterization in ALL

Array-based technologies

The advent of microarray-based technologies provided a leap from low- to high-throughput analysis that transformed the settings for molecular biology studies [46]. Microarray technologies comprise numerous applications including genome-wide profiling of single nucleotide polymorphisms (SNP), copy number alterations (CNA), gene expression patterns, and transcription factor binding site profiling [47]. Although microarrays are limited in their genomic coverage compared to NGS technologies, they are particularly advantageous for large-scale studies involving numerous samples and provide
a straightforward approach for analyzing hundreds of thousands of genomic sites at a reasonable cost.

**SNP and gene expression arrays in ALL**

Since 2007, SNP arrays have been used to uncover a spectrum of secondary CNAs (DNA deletions or gains), that are important cooperating lesions for disease pathogenesis, especially in BCP-ALL [48-50] Transcription factors that are essential for hematopoiesis, including *PAX5*, *IKZF1*, *ETV6* and *EBF1* are among the most frequently altered genes in addition to the tumor suppressors and cell cycle regulators *CDKN2A/B*, *PTEN* and *RB1* [51-53]. In fact, the crucial B-cell lineage regulating transcription factor *PAX5* is recurrently mutated, deleted or rearranged in over one-third of BCP-ALL cases [52]. The type and frequency of *PAX5* alterations vary between the ALL subtypes, and *PAX5* deletions occur in up to 36% of B-other cases [54]. Deletions of the lymphoid transcription factor *IKZF1* are reported in 15% of BCP-ALL cases, but is more common in the t(9;22)*BCR-ABL1* subtype where *IKZF1* deletions are found in >60% of the patients [55, 56]. Unlike the gross chromosomal rearrangements and aneuploidies that are usually initiating lesions acquired early during the leukemic transformation, the submicroscopic DNA changes represent subsequently acquired alterations [51, 57-60]. Some of the CNAs, such as deletions of *PAX5*, *IKZF1* or *CDKN2A/B*, have been shown to promote the onset of leukemia in mouse models, supporting their role as central cooperative lesions for leukemogenesis [61-63].

Before the advent of NGS, gene expression arrays represented a unique platform to study differential expression of a defined set of target genes. Early microarray-based expression studies in ALL could establish that the different cytogenetic subtypes of ALL have unique gene expression signatures, which led to a considerable better molecular understanding of the subtypes of ALL [64-66]. In 2009, a new subtype termed *BCR-ABL1*-like (also known as Ph-like) was described in the B-other group with the aid of gene expression microarrays [56, 67]. This group of patients was characterized by an expression profile similar to that of the t(9;22)*BCR-ABL1* subtype, yet they lack the canonical *BCR-ABL1* fusion gene detectable by gene expression arrays. Similarly to the t(9;22)*BCR-ABL1*, the *BCR-ABL1*-like subtype also harbors high frequencies of *IKZF1* deletions, and are associated with a poor outcome [56, 67, 68]. Today, *BCR-ABL1*-like is recognized as a new provisional entity of leukemia that comprises 10-15% of the ALL patients and is classified as a high-risk marker [69].

**RNA-sequencing**

The advancements in NGS techniques, coupled with the reduction in sequencing costs, have enabled new opportunities for comprehensive analysis of the transcriptome. Understanding the abundance and structure of RNA is
essential for interpreting molecular processes involved in maintaining cell functions and for disease pathogenesis. For this purpose, RNA-sequencing offers unbiased determination of gene expression levels, expressed variants, alternative splicing and detection of fusion genes. Multiple protocols for RNA-sequencing exist, typically involving enrichment or depletion of “total” RNA for specific RNA molecules, i.e. polyadenylated RNA and ribosomal RNA, respectively, followed by conversion of the remaining RNA of interest to cDNA prior to sequencing library preparation [70].

**RNA-sequencing in ALL**

RNA-sequencing is a powerful tool for detection of known and novel fusion genes. In ALL, RNA-sequencing it is particularly advantageous due to the high frequency of expressed fusion genes as a consequence of recurrent translocations. In light of this, recent studies taking advantage of RNA-sequencing in BCP-ALL has uncovered the presence of several key fusion genes in the B-other subtype that are associated with clinical outcome and are thus of potential clinical relevance. The most notable findings have been the recurrent fusion gene rearrangements involving DUX4, ZNF384 and MEF2D [32, 44, 45].

The DUX4-rearranged subtype is characterized by a fusion gene resulting from the insertion of the DUX4 gene into the IGH locus on chr14 and recurrent ERG deletions [71-74]. Before the discovery of expressed DUX4-related fusion genes, prior studies had reported recurrent deletions of the ERG gene in approximately 5% of ALL patients and this putative subgroup of ALL patients was thus referred to as “ERG-deleted” [75-77]. Today, it has been established that ERG deletions are restricted to the DUX4-rearranged subtype. The DUX4-rearranged subtype has so far been reported in up to 7% of the BCP-ALL cases and appears to have a favorable prognosis under current clinical protocols in multiple studies [44, 71, 72, 78].

Rearrangements involving ZNF384 and MEF2D are less frequent, with estimated frequencies of 1-6% and 1-4% in BCP-ALL, respectively [32, 45]. Both ZNF384 and MEF2D rearrangements have been reported to involve multiple gene fusion partners [72, 73, 79-82]. Patients with ZNF384 have been reported to have varying clinical outcome depending on its 3’ fusion partner (denoted as intermediate prognosis), whilst cases harboring MEF2D rearrangements are associated with poor clinical outcome [45, 79]. Due to the small sample sizes analyzed to date, larger studies will be required in order to fully establish the prognostic relevance of these two novel subtypes.

The BCR-ABL1-like subtype, as described above, is a heterogeneous subgroup involving different genetic alterations that activate kinase and cytokine receptor signaling and is associated with a dismal outcome [68]. Apart from its aforementioned molecular resemblance to the t(9;22)BCR-ABL1 subtype, RNA-sequencing has revealed a plethora of expressed fusion gene within the BCR-ABL1-like subtype, involving tyrosine kinase receptor genes
such as PDGFRB, ABL1, ABL2 or CSF1R, and JAK/STAT activating rearrangements involving CRLF2, JAK2 and EPOR [83-86]. A subset of the BCR-ABL1-like patients lack these rearrangements, but harbor other less frequent JAK/STAT activating lesions or rare kinase related fusion genes [68]. Gene expression profiling is helpful for discerning B-other patients with putative BCR-ABL1-like expression patterns and a common approach relies on microarray-based gene expression signatures from a predefined probe set [87, 88]. For cases where gene-expression microarray data is not available, RNA-sequencing has also been successfully applied for identification of these patients based on gene expression signatures that are similar to t(9;22)BCR-ABL1 patients [71, 73]. Importantly, as a consequence of lesions involving tyrosine kinases, the use of tyrosine kinase inhibitors could improve outcome for the patients in this subgroup [84, 89].

Whole genome sequencing

Whole genome sequencing (WGS) is a powerful technique for high-resolution profiling of genetic variation and has revealed numerous somatic single nucleotide variants (SNV) and mutational signatures that are important contributors for cancer progression. By genome-wide profiling, WGS enables de-novo detection of somatically acquired mutations in non-coding regions including introns and regulatory regions such as enhancers in an unbiased manner [90]. Despite its strength, WGS is limited in identifying the full spectrum of structural variants such as translocations, insertions and deletions [90]. Furthermore, complex genomic regions such as repetitive sequences are still challenging to fully resolve in detail by standard short-read WGS data.

Recently, a new technology termed “linked-read” WGS (10x Genomics) was introduced [91]. Linked-read WGS enables improved resolution across complex genomic regions and detection of structural variants, owing to its capacity to generate the long-range phased genomic regions that is beyond the capacity of standard short-read sequencing. The principle of long-read sequencing is to utilize massive barcoding strategies and microfluidics for partitioning single DNA molecules to generate barcoded sequencing libraries that are subjected to standard short-read WGS. After WGS the sequencing reads are aligned and assembled into synthetic “long-read” genome sequences with the aid of unique barcodes for the WGS reads generated from the same DNA molecule. This enables computational reconstruction of long haplotype blocks and thus phasing of genetic variants including, chromosomal rearrangements, single nucleotide variants, and small insertion and deletions (Figure 3). Linked-read WGS offers a promising approach for assignment of complex variants, such as SNVs, insertion-deletions, chromosome breaks, gene fusions and other rearrangements to individual chromosomes in germline and cancer genomes [91-93]
Figure 3. Reconstruction of phased blocks by linked-read whole genome sequencing. The colored circles represent short-read sequences that are linked together based on their common barcodes to generate unambiguous haplotype blocks for the homologous chromosome pairs. A deletion on haplotype 2 is illustrated by the absence of short reads aligning to that region.

WGS in ALL

The mutational landscape of ALL has been extensively studied with the aid of high-throughput sequencing such as short-read WGS [51]. Opposite to gene deletions, the frequency of somatic mutations is low in ALL, and the 11q23/MLL subtype has been found to have amongst the lowest mutation rates of all cancers [94, 95]. Indeed, a spectrum of mutations has been detected in ALL, including mutations in members of the RAS signaling pathway (NRAS, KRAS) and hypodiploid-associated TP53 mutations [96-99]. The most notable findings has been generated by studies of the clonal evolution in relapsed ALL that have identified genes that are enriched at relapse such as activating NT5C2, that drives chemotherapy resistance in ALL and is detected in up to 20% of relapsed cases [100, 101].

Characterization of the DNA methylome in ALL

DNA methylation is an epigenetic modification of the DNA that occurs predominantly at CpG dinucleotides (a cytosine followed by a guanine), which is mediated by DNA methyltransferases (DNMTs). The modification involves the addition of a methyl group to cytosine; giving rise to a “5th base” i.e. 5-methylcytosine. DNA methylation is a robust and stable epigenetic mark that can been faithfully analyzed with various techniques, including microarray-based genotyping, sequencing-based approaches and mass-spectrometry [102].

Infinium microarrays provide a straightforward assessment of the DNA methylation status of many thousands of CpG sites in large sample sets, at a relatively low cost. Infinium microarrays for DNA methylation analysis include predetermined CpG sites distributed across the genome, which cover 99% of the genes in the Reference Sequence database [103]. The number of target probes included on the arrays have increased over the years, starting
with the HumanMethylation 27k array (targeting ~27000 CpG sites), fol-
lowed by the 450k array and the EPIC array (targeting over 850 000 CpG
sites) that covers both genes and enhancer regions [104-106]. It is to be not-
oted that DNA methylation arrays target only up to 3% of the 28 million CpG
sites present in the human genome and alternative methods such as whole-
genome bisulfite sequencing are required for complete genome-wide DNA
methylome assessment at single-base pair resolution [107].

In healthy cells, it has been estimated that 60-80% of the 28 million CpG
sites present in the human genome can become methylated. Unmethylated
CpGs are typically clustered together in about 200 bp long CpG-rich regions
(“CpG islands”) and are enriched in gene promoters [108, 109]. DNA meth-
ylation plays an important role in orchestrating cell-type specific gene ex-
pression, giving each cell type its specific identity. In general, but not al-
ways, a high DNA methylation level (hypermethylation) in promoter regions
silences gene expression [108].

DNA methylation in ALL

In ALL, the genome-wide DNA methylation landscape is altered and the
methylation pattern of the ALL cells are clearly distinct from that of normal
healthy blood cells [64, 110, 111]. Changes in DNA methylation in cancer
cells are mainly seen as hypermethylated CpG islands compared to normal
controls and has been observed across all genetic and immunophenotypic
subtypes of ALL [112]. These differentially methylated CpG sites (DMCs)
that are shared between the genetic subtypes of ALL suggests a common
mechanism for leukemic transformation in ALL. Further dissection of the
DNA methylation landscape across the different subtypes of BCP-ALL has
identified strong subtype-specific DMCs that are enriched in promoter and
enhancer regions [64, 110-112].

In the context of ALL prognosis, it has been shown that promotor-
associated DNA methylation holds prognostic information that can predict
relapse in T-ALL [113]. In this study, patients were stratified based on a
CpG island methylator phenotype (CIMP) profile, where patients with lower
DNA methylation levels at diagnosis (CIMP-) were more likely to relapse,
compared to patients with high methylation levels (CIMP+) where relapses
were less frequent. Recently, it was shown that CIMP classification is also
important for relapsed BCP-ALL and correlates with overall survival after
relapse [114].

DNA methylation classification in ALL

Today, it is well-acknowledged that each genetic subtype of BCP-ALL is
associated with a distinct set of differentially methylated CpG sites that can
be used to distinguish between the subtypes [110]. This feature can be uti-
liized to design DNA methylation-based classifiers to predict the cytogenetic
subtype of an ALL sample, solely based on DNA methylation status of a
limited set of CpG sites [115]. Classifiers for the most common and established subtypes of BCP-ALL (HeH, t(12;21)ETV6-RUNX1, t(1;19)TCF3-PBX1, 11q23/MLL, t(9;22)BCR-ABL1, dic(9;20) and iAMP21) in addition to T-ALL, have been developed [115]. Interestingly, when the DNA methylation classifiers were applied to a large set of previously undefined B-other samples, nearly half of the samples displayed highly similar DNA methylation patterns as samples with a known subtype at diagnosis. Verification of the predicted subtype in B-other cases led to the discovery of several unexpected fusion genes, including a small number of patients with similar DNA methylation profiles as positive t(12;21)ETV6-RUNX1 cases, however alternative fusion genes involving either ETV6 or RUNX1 were detected, instead of the canonical ETV6-RUNX1. Recently, additional patients with ETV6-RUNX1-like profiles have been reported based on gene expression or immunophenotypic patterns that are similar to positive t(12;21)ETV6-RUNX1 cases, and are now referred to as the “ETV6-RUNX1-like” group [71, 116]. ETV6-RUNX1-like cases frequently harbor co-occurring fusion genes and/or deletions involving ETV6 and IKZF1 [44, 71, 116]. Very few relapses have been reported in the ETV6-RUNX1-like subtype, suggesting a good prognostic outcome for this putative subtype, similarly to the canonical t(12;21)ETV6-RUNX1 subtype. Larger studies are required to fully discern the prognostic significance of this emerging subtype.

Not restricted to ALL, DNA methylation as a marker for classifying cancers has been successfully applied to a wide spectrum of malignancies to discriminate between different molecular and outcome groups [117-120]. Together, this reflects a potential use of DNA methylation as a future biomarker.
Present investigations

Thesis aims
The overall aim of this thesis was to study genetic alterations in pediatric acute lymphoblastic leukemia (ALL) patients using multiple genome-wide techniques. More specifically the aims were to:

- Decipher the genetic alterations in patients with ALL focusing on patients lacking known risk-stratifying markers (B-other).
- Explore novel genomic strategies for detection of ALL subtypes based on in-depth analyses of structural rearrangements and fusion genes.
- Gain further understanding of the frequency and prognostic relevance of the identified genetic alterations.

Materials and methods
Several methods for genome-wide analysis were applied throughout the work presented in this thesis. The following section highlights the most important methods used in this work and detailed information about each of these different methods is presented in the individual papers I-IV.

Patient samples and clinical data
The samples from patients with pediatric ALL analyzed in this thesis belong to a unique collection of bone marrow and peripheral blood samples obtained from >1000 children diagnosed with ALL during 1996-2012 within the Nordic countries. The samples were collected at collaborating institutes within the Nordic Society of Pediatric Hematology and Oncology (NOPHO). Phenotypic data has been thoroughly recorded since sample collection in the NOPHO patient registry, which enables follow-up analysis the clinical outcome of the patients over an extended period of time.

Primary leukemic cells collected at initial diagnosis were isolated by Ficoll-Paque centrifugation prior to cryopreservation. Extraction of DNA and RNA was performed using column-based extraction methods including a
DNase treatment step of the RNA. High molecular weight DNA (HMW DNA) was extracted from two samples (Paper IV). Isolated material was quantified using fluorescent assays and quality of the RNA was assessed using the RNA integrity number (RIN) algorithm which is based on the ratio of measured ribosomal RNA 28S and 18S.

Clinical outcome of the patients was assessed by calculating the probabilities of event free survival and overall survival using the Kaplan-Meier method. For event-free survival estimation, an event was defined as resistant disease, induction failure, relapse, death in remission or second malignant neoplasm.

DNA methylation analysis
For genome-wide interrogation of the DNA methylation levels of >450,000 CpG sites genome-wide in the leukemic samples, the Human Methylation 450k BeadChip (450k array, Illumina) was utilized. Sodium bisulfite treatment was performed on 250-500 ng of genomic DNA using the EZ DNA Methylation-Gold Kit (Zymo Research). The bisulfite step converts unmethylated cytosines (C) to uracils, which become thymines (T) after PCR, whilst methylated cytosines remain unaffected. By genotyping the C/T polymorphisms using fluorescent probes, the methylation levels across the predefined CpG sites included on the 450k array can be determined.

Raw fluorescent signal intensities from the 450k array were detected and quality controlled with the Minfi R package [121]. The level of DNA methylation for each CpG site was transformed into β (beta) values, which corresponds to the intensity signal from the methylated C-nucleotides over the sum of the signals from the C- and T-nucleotides using GenomeStudio (Illumina). The β-values range from 0-1 which corresponds to 0-100% methylation for each CpG site included on the 450k array. After normalization of the β-values using the peak-based correction algorithm [122], probes with a detection p-value >0.01 or with SNPs or indels with minor allele frequencies >5% within the last 3 bp of the 3’end of the primers were excluded from downstream analyses [110].

RNA-sequencing and fusion gene detection
Strand-specific next generation RNA-sequencing libraries were constructed from 100-300ng total RNA following ribosomal RNA depletion. High-quality RNA samples (RIN values >7) were used for library preparation using the ScriptSeq V2 Kit (EpiCentre) or the TruSeq stranded Total RNA kit (Illumina). A subset of the sequencing libraries from RNA samples of lower quality (RIN <7) were prepared using the TruSeq RNA Access kit (Illumina). The libraries were sequenced using Illumina systems (HiSeq2000, 2500, Miseq or NovaSeq) with 50-150bp paired-end chemistry, generating
20-40 million read pairs per sample. In paper III, Cutadapt [123] was used to trim the sequencing reads to 65 bp to obtain a uniform sample set as different sequencing platforms had been used. Sequencing reads were quality controlled using FastQC [124] prior to alignment against the human reference genome (hg19) with the TopHat2 or STAR software [125, 126]. RNA-sequencing data was quality-controlled using either the RNA-SeQC or RSeQC software [127, 128].

Fusion genes were detected using the FusionCatcher software [129]. Targeted screening of several well-established fusion genes in ALL was performed by counting reads supporting fusion transcripts in the TopHat2-aligned data normalized to the total sequencing depth in the sample. A multi-step filtering process of the candidate fusion genes detected by FusionCatcher was applied to reduce the number of false positives. Briefly, these steps included filtering of blacklisted fusion genes based on reference data provided by FusionCatcher followed by removal of fusion genes based on; (i) common mapping of fusion-supporting reads to multiple positions, which is indicative of sequence homology; (ii) highly promiscuous genes where the 5’ or the 3’-fusion gene partner maps to several other genes; (iii) few sequencing reads supporting the fusion junction; or, (iv) expression of the fusion gene in normal CD19+ B-cells and CD3+ T-cells from healthy blood donors.

Gene expression profiling

Quantification of gene expression levels from the RNA-sequencing data was performed using Cufflinks (version 2.2.0) or featureCounts (Subread version 1.5.0) [130, 131]. FeatureCounts was used to summarize the counts of aligned reads and normalized using variance stabilizing transformation with DESeq2 [132]. Differentially expressed genes were analyzed within the Cufflinks pipeline by determining the fragments per kilobase per million mapped reads (FPKM) levels. Cuffdiff was subsequently used to detect differentially expressed genes by pair-wise comparison of the expression levels between two ALL subgroups [133]. Selection of differentially expressed genes in paper III was based on a false discovery rate corrected p-value <0.1 and a two-fold difference in mean expression detected between three or more compared ALL groups. Genes with low expression levels (FPKM <5) were excluded from the analysis.

Copy number analysis

Illumina Infinium microarrays were used to screen for copy number alterations. Several Illumina Infinium arrays were utilized, including the 450k array for DNA-methylation analysis, Omni2.5 array, OmniExpressExome array and the Multi-Ethnic Global array.
Notably, the 450k array is based on the same biochemical principles as the regular SNP genotyping arrays, i.e. using two fluorophores to interrogate the loci included on the bead arrays. Thus, the total raw fluorescence intensity from the 450k array can also be used to detect copy number variants in genomic DNA. The raw signal intensities generated from the 450k array were normalized and transformed to “Log R ratio”, which corresponds to the observed intensity signal from the studied sample over the intensity signal obtained from a reference sample (baseline). To determine the reference baseline, 450k data from CD19+ B-cells and CD3+ T-cells from healthy blood donors representing normal diploid genomes, was used. Log R ratios were analyzed using the CopyNumber450kCancer R package [134].

Log R ratios from the high-density SNP arrays were calculated in GenomeStudio (Illumina) to assess copy number status using an in-house panel of diploid samples or with Illumina standards for reference intensities based on HapMap samples. In addition to analyzing Log R ratios for copy number status, genotyping of A/C/T/G polymorphisms for detection of copy neutral allelic imbalances by estimation of the B-allele frequency was performed in GenomeStudio. Copy number alterations were called from the SNP array data using the Tumor Aberration Prediction Suite (TAPS) analysis software [135].

**Linked-read whole genome sequencing**

Linked-read whole genome sequencing (linked-read WGS) technology for haplotyping (phasing) of genetic variations was evaluated in study IV. For this purpose, the GemCode and/or the Chromium technology from 10x Genomics was applied. This technology uses a combination of microfluidics and molecular barcoding to generate WGS libraries [91]. The resulting linked-read sequencing libraries enable computational reconstruction of haplotype and structural variant information within a single WGS experiment. Briefly, long DNA molecules are partitioned into single oil drops together with gel beads loaded with enzymes, primers and barcodes. As little as 1 ng of genomic DNA is sufficient as starting material, which is equivalent to approximately 300 copies of the human genome. The resulting barcoded DNA fragments are subsequently sheared, pooled, and finally a standard Illumina short-read sequencing library is generated. GemCode libraries were sequenced on an Illumina HiSeq2500 and Chromium libraries were sequenced on the HiSeqX instrument. A customized sequencing protocol was used for the GemCode libraries whereas standard 150bp paired-end chemistry was used for the Chromium libraries. The linked-read data was processed using a DNA analysis pipeline available from 10x Genomics (Long Ranger) that performs sample de-multiplexing, barcode processing, alignment, quality control and phasing. Structural variant calling was performed using Long
Ranger and CNVnator [136]. Identified structural variants by Long Ranger were visualized in the Loupe genome browser (10x Genomics).

Results

Identification of the PAX5-ESRRB fusion gene (Study I)

In study I, we describe a novel and recurrent fusion gene using a combination of RNA-sequencing, reverse transcription PCR (RT-PCR) and copy number detection in B-other. These patients where characterized by the expression of PAX5-ESRRB, a fusion gene stemming from an unbalanced translocation between chr9 and chr14 (Figure 4). This fusion gene remained undetected at diagnosis using traditional karyotyping in each of the four patients. Owing to the unbalanced nature of the translocation, we utilized the Log R ratio from the 450k array data to screen for copy number alterations involving chr9 and chr14 in 664 pediatric BCP-ALL cases. These patients had previously been analyzed using the 450k DNA methylation array, and the frequency of this fusion was estimated to be <1% in BCP-ALL (4/664). The four ALL cases carrying the PAX5-ESRRB fusion gene were at clinical remission after follow-up for 30-173 months. At a molecular level, patients carrying the PAX5-ESRRB fusion gene showed distinct gene expression patterns characterized by high expression of ESRRB, in addition to a predominant hypomethylation of DNA, as compared to other BCP-ALL subtypes.

Figure 4. Chromosomal rearrangements giving rise to the expressed PAX5-ESRRB fusion gene. (A) FISH analysis revealed the presence of a der(9)t(9;14) rearrangement resulting from a translocation between chromosome 14q to chromosome 9p. (B) Sanger sequencing revealed the expression of an in-frame PAX5-ESRRB fusion gene in all cases. Exons involved in the fusion transcript are represented by the numbered squares in the upper panels and color coded to indicate the specific genes involved. Sanger sequencing traces spanning across the fusion junction (vertical dashed line) are illustrated in the lower panel.
Fusion genes with distinct DNA methylation profiles (Study II)

In study II, we further explored the fusion gene landscape of ALL in an unbiased manner by performing total RNA-sequencing on 134 ALL patient samples assigned to various subgroups, including B-other. After rigorous filtering of candidate fusion genes called by FusionCatcher, a total of 64 unique fusion genes were identified in 80 of the 134 (60%) patients included in the study. Subtype-defining fusion genes including ETV6-RUNX1, BCR-ABL1 and KMT2A-chimeras, were unambiguously detected in 29 of the 31 BCP-ALL cases with the established subtypes t(12;21), t(9;22) or 11q23/MLL.

Although most fusion genes (43/64; 67%) were only detected in a single patient, we detected several previously undetected and recurrent fusion gene families in over 80% (35/42) of the patients in the B-other group. Several of these fusion genes arose due to cryptic or subtle structural rearrangements that were not detectable by traditional diagnostic methods. The most recurrent fusion genes that were detected in 50% of the analyzed B-other transcriptomes, involved the DUX4, ZNF384 or PAX5 genes. Whereas multiple gene partners were found to fuse with the ZNF384 and PAX5 genes, DUX4 fusion genes involved the IGH locus located on chromosome 14 in each case. Furthermore, similar to the well-established subtypes of ALL, the putative subtypes harboring DUX4 and ZNF384 displayed distinct subtype-specific DNA methylation and gene expression profiles (Figure 5). Using rapid amplification of cDNA ends (RACE), two additional patients harboring DUX4 and ZNF384 fusion genes were identified. These patients were discovered based on similar DNA methylation and gene expression profiles as the verified DUX4-rearranged and ZNF384-rearranged cases, however our screening strategy was not able to detect a fusion gene involving DUX4 or ZNF384 in the RNA-sequencing data. Further characterization of the DUX4-rearranged and ZNF384-rearranged cases revealed a high prevalence of NRAS mutation in the DUX4 subgroup (56%) in addition to frequent ERG deletions (78%). Assessment of prognostic relevance for patients with DUX4 fusion genes confirmed a good prognosis, and an intermediate prognosis in patients harboring the ZNF384 fusion genes, as recently reported by other studies [72, 73, 79].

Amongst the sequenced B-other cases, three patients previously described to have identical DNA methylation patterns and classified to have the t(12;21)ETV6-RUNX1 subtype were included in this study. These patients had previously been found to harbor novel, alternative ETV6 or RUNX1 fusion genes instead of the canonical ETV6-RUNX1 fusion gene [115]. In agreement with the DNA methylation profile, these three patients also clustered with t(12;21)ETV6-RUNX1 samples based on their gene expression profiles.
DNA methylation classifiers for novel ALL subtypes (Study III)

In study III, subtype-specific DNA methylation patterns in the common subtypes of ALL were utilized [115], together with new insights on the putative DUX4 and ZNF384 subtypes reported in study II, to design DNA methylation-based classifiers for ALL. For this purpose, we further refined our previously designed supervised learning procedure based on nearest shrunken centroids, to expand our original DNA methylation classifier by inclusion of the novel subtypes reported in study II. Classifiers were built for 10 ALL subtypes: T-ALL, HeH, t(12;21)ETV6-RUNX1, t(1;19)TCF3-PBX1, 11q23/MLL-rearrangements, t(9;22)BCR-ABL1, dic(9;20), iAMP21, DUX4- and ZNF384-rearrangements.

An external cohort of 243 blinded ALL samples with an established subtype at ALL diagnosis, was used to assess the performance of the classifier, which resulted in unequivocally assignment of 87% of the ALL cases to the correct subtype. For validation of the subset of samples for which the classifier failed to assign the correct subtype (n=31), a lower than expected blast count was observed, which could have confused the classifier and account for the discordant results. One sample diagnosed as HeH was classified as a DUX4-rearranged subtype. This finding was validated by RNA-sequencing, which confirmed the expression of the DUX4-IGH fusion gene.

Next, the designed classifiers were applied as a tool to screen for subtype membership in 284 B-other cases. Over 50% of the B-other patients were assigned to an existing ALL subtype using this DNA methylation-based classification method (Figure 6). Interestingly, the most commonly predict-
ed subtype was the *DUX4*-rearranged subtype which together with the *ZNF384*-rearranged subtype accounted for 15% (10% and 5%, respectively) of the B-other samples. To confirm the predicted subtypes, based on their DNA methylation signature, copy number analysis was performed in addition to RNA-sequencing for cases with RNA available. RNA-sequencing identified several novel and known fusion genes in the B-other patients, which confirmed the new subtype class including *DUX4*-rearranged, *ZNF384*-rearranged, *BCR-ABL1*-like and *ETV6-RUNX1*-like.

The emerging *ETV6-RUNX1*-like subtype was characterized fusion genes involving the *ETV6* or *IKZF1* genes together with frequent co-occurring *ETV6* deletions, and with no detectable expression of the canonical *ETV6-RUNX1* fusion gene. Of note, several of these samples had in fact been negative for t(12;21)*ETV6-RUNX1* by FISH or PCR at diagnosis.

![Figure 6. DNA methylation classification in B-other. Distribution of predicted subtypes in B-other including samples that were assigned to more than one class (multi-class) and samples that did not get a subtype score (non-class) for any of the subtypes included in the classifier.](image)

**Digital karyotyping by linked-read WGS (Study IV)**

In study IV, we evaluated novel linked-read WGS that enables increased resolution compared to standard WGS and phasing of structural variants on each chromosome. Linked-read sequencing was applied for analysis of DNA samples from 12 ALL patients and one ALL cell line (REH) of varying quality. To ensure optimal phasing and computational reconstruction of long haplotypes, isolation of high-molecular weight DNA (HMW DNA) is crucial. We assessed the effect of the quality of input DNA by comparing HMW DNA to that of DNA extracted with standard column-based DNA preparations and had undergone repeated freeze-thawing without selection for long DNA fragments. Although shorter phasing blocks were assigned using standard DNA samples for linked-read WGS, comparable results were
obtained for detection of structural variants in two control samples harboring the \( t(12;21)\)ETV6-RUNX1 and \( t(9;22)\)BCR-ABL1 fusion genes.

Linked-read WGS enabled precise digital karyotyping of a wide range of structural variants such as balanced and unbalanced translocations, deletions and aneuploidies. Complex rearrangements involving up to five different chromosomes that were either missed or incorrectly annotated by traditional karyotype analysis were resolved at single-base pair resolution by linked-read WGS (Figure 7). As a proof of principle, digital karyotyping detected all subtype-defining events reported at diagnosis in the HeH, \( t(12;21) \) and \( t(9;22) \) subtypes analyzed in this study. Co-occurring \( ETV6 \) deletions were found in all \( t(12;21)ETV6-RUNX1 \) patients and by utilizing phasing information, we were able to better resolve the deletion of \( ETV6 \) on the allele that was not affected by the translocation in one patient.

Furthermore, the spectrum of genetic alterations in B-other patients harboring fusion genes involving \( DUX4 \), \( ZNF384 \) or \( PAX5 \) were delineated at a resolution beyond that of traditional karyotyping or short-read WGS. Owing to the phasing ability of the technology, a previously unobserved bi-allelic \( ERG \) deletion was discovered in a patient with the \( DUX4-IGH \) fusion gene (Figure 7). Furthermore, we detected heterozygous \( IKZF1 \) deletions in both \( DUX4-IGH \) cases subjected to linked-read WGS.

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**Figure 7.** Genomic view of structural rearrangements in ALL by linked-read WGS. (A) Digital karyotyping by linked-read WGS resolved a complex translocations involving five chromosomes (chr 2, 3, 12, 14 and 21) in addition to the subtype defining \( t(12;21)ETV6-RUNX1 \) aberration indicated with an arrow. (B) Genomic view of the linked-reads mapped to each of the two chromosomes at the \( ERG \) locus on chr21 in a BCP-ALL patient with the \( DUX4-IGH \) fusion gene. Reads are color-coded by chromosome (haplotype 1 in green and haplotype 2 in purple) and deletions are marked by red squares.
Summarized discussion

B-other represents a large subgroup of ALL patients where the chromosomal aberrations remain largely undescribed based on currently used diagnostic methods. Identification of the key alterations that define new oncogenic subtypes within the B-other group is imperative, as these patients may benefit from alternative treatment regimens such as more or less intensive treatment protocols or in some cases addition of targeted therapies.

The studies summarized in this thesis have together resolved several recurrent alterations in the B-other group using different genome-wide techniques for molecular profiling in >1000 pediatric ALL patients that were diagnosed and treated according to NOPHO protocols. The large collection of genome-wide DNA methylation data represents a unique backbone for this thesis, while RNA-sequencing and linked-read WGS performed on a subset of patients supports that modern technologies improves the resolution of the chromosomal alterations, especially in the B-other subgroup, by offering precise base-pair delineation of the genetic alterations. Due to the unique nature of the ALL cohort described in this study and the availability of clinical and follow-up data for these patients, we have shown that several of the newly identified alterations detected in B-other [32, 44, 45] are recurrent and prognostically relevant in ALL.

One of the most recurrent alterations discovered in the B-other group is the $\text{DUX4-IGH}$ fusion gene, which we estimate to be present in approximately 4% of BCP-ALL (14% in B-other). Similar to what has been seen in other studies, NOPHO BCP-ALL patients with $\text{DUX4}$-rearrangements have a favorable outcome and $\text{ERG}$ deletions are frequent in this group. Owing to the phasing ability of linked-read WGS we were able to better resolve a submicroscopic $\text{ERG}$ deletion in one $\text{DUX4-IGH}$ patient and identified a compound heterozygous $\text{ERG}$ deletion, that could not be resolved by RNA-sequencing or copy number analysis, and was undetected in a previous short-read WGS study where this patient was sequenced [98].

Opposite to the $\text{DUX4}$-rearranged subtype, $\text{BCR-ABL1}$-like is considered a high-risk subtype with dismal prognosis and comprises various kinase-activating lesions [87]. The NOPHO patients harboring tyrosine kinase fusion genes identified by RNA-sequencing in the present studies suffered from frequent relapse and fatal outcomes. Thus, these results strengthen pre-
vious observations that patients with the *BCR-ABL1*-like phenotype may benefit from alternative treatment strategies, such as the addition of targeted tyrosine kinase inhibitors [84].

Moreover, this is the largest study addressing the putative *ETV6-RUNX1*-like subtype to date, in which we confirm the favorable prognosis associated with this emerging subtype and frequent co-occurring *ETV6* deletions [71, 116]. Our findings strengthens that *ETV6-RUNX1*-like is a recurrent putative subtype in BCP-ALL and comprises 1.4-2% of the cases (5-7% in B-other). Additional functional studies will be required in order to understand the molecular mechanism for the *ETV6-RUNX1*-like subtype and its molecular and clinical similarity to the classical t(12;21)*ETV6-RUNX1* subtype, although one might speculate that alterations of the *ETV6* gene is a common denominator between these analogous subtypes.

Interestingly, co-occurring *ETV6* deletions are also common in positive t(12;21)*ETV6-RUNX1* patients and detected in up to 60% of the cases [54, 137]. In each of the three t(12;21)*ETV6-RUNX1* patients we analyzed by linked-read WGS, each case harbored a co-occurring *ETV6* deletion on the non-translocated allele, resulting in a bi-allelic disruption of *ETV6*. Using haplotype information from the linked-read WGS data, the entire *ETV6* locus was phased in one t(12;21)*ETV6-RUNX1* patient that resolved an intergenic *ETV6* deletion on one allele and the translocation forming the *ETV6-RUNX1* fusion gene originating from the other allele.

Fusion genes involving *PAX5* occur in approximately 2.5% of BCP-ALL and is a known heterogeneous fusion gene partner [138]. In our RNA-sequencing studies, we found that *PAX5* forms chimeric transcripts with up to 10 different genes in the patients included herein. Together, fusion genes involving *PAX5* were detected in approximately 2% of the BCP-ALL cases and is thus in line with previous findings [138].

Unlike the established and new putative subtypes of ALL such as those with *DUX4*- and *ZNF384*-rearrangements, the heterogeneous group of patients with *PAX5* fusion genes does not appear to have a strong and distinctive DNA methylation or gene expression pattern (Figure 5). Functional studies of *PAX5* chimeras have established both common and distinct properties associated with the different fusion gene partners that fuse to *PAX5* [139]. Notably, the different fusion gene partners may affect the DNA-binding affinity and regulatory potential of the *PAX5* fusion gene in repressing or activating its target genes, consequently affecting the molecular features of the leukemic cells [139].

In particular, we described for the first time a recurrent subgroup characterized by the *PAX5-ESRRB* fusion gene from an original cohort of 664 Nordic BCP-ALL samples diagnosed before 2008. The four patients with *PAX5-ESRRB* in our study did display distinct DNA methylation profiles that dif-
ferred from that of other \textit{PAX5} fusion genes, suggesting that there is some degree of unique molecular signatures, at least in this small group of patients. Nevertheless, this analysis was performed in a small sample set and warrants further investigation in order to understand the diversity of \textit{PAX5} chimeras and to establish whether other recurrent \textit{PAX5} chimeras share distinct and unique molecular features. Interestingly, although we performed copy number analysis and RNA-sequencing in additional recently diagnosed B-other patients, no other patients with \textit{PAX5-ESRRB} were detected in our studies nor have other studies on BCP-ALL reported this \textit{PAX5} fusion gene, which suggests that this fusion gene is extremely rare in BCP-ALL.

Although the studies presented in this thesis resolved the genomic complexity for over half of the patients in the B-other group thus enabling an improved delineation of the genetic alterations in this subgroup, the underlying genetic alteration still remains unknown for a subset of the patients. Consequently, further analysis is required for these patients in order to fully resolve the lesions driving the leukemia, and potentially identifying new subtypes.

From a technical perspective, we demonstrate the power of DNA methylation as a marker for ALL subtype prediction that can be used to retrospectively revise the subtype membership of a large proportion of previously undefined B-other cases. A main strength with this approach is that subtype membership can be given to patients diagnosed over 20 years ago. This reflects the stability of DNA methylation as a biomarker even for DNA samples that were extracted from cryopreserved patient samples that had been biobanked for a long time. Nonetheless, the DNA methylation profile alone is currently not sufficient to confidently establish a subtype membership, and we must still rely on complementary methods such as copy number analysis or verification of expressed fusion genes by RT-PCR, RNA-sequencing or RACE, to unequivocally assign a new subtype to an ALL patient.

We found that our DNA methylation classifiers are sensitive to leukemic cell composition. Samples with lower leukemic blast count could not be accurately assigned to the expected subtype, whereas RNA-sequencing from the same sample with low blast count successfully identified the expected fusion gene. The restricted dynamic range of detection by DNA methylation as compared to RNA-sequencing, which offers a greater dynamic range and is less sensitive to tumor content, is an important aspect when considering a robust test for clinical diagnostics.

Although DNA methylation classification has been a valuable tool for screening of ALL subtypes in the large retrospective cohort described in this thesis, it is unlikely that this approach can be implemented as a diagnostic test for ALL in a clinical setting. Instead, methods that are based on fusion gene detection, such as RNA-sequencing or similar, would be more conven-
ient as they provide a direct answer of expressed fusion transcripts and its involved gene partners. Knowing the exact fusion gene may be beneficial for MRD screening [41] and also for subtypes such as ZNF384-rearranged and 11q23/MLL as they may confer different clinical outcomes depending on the fusion gene partner [79, 140].

Lastly, we evaluated the utility of linked-read WGS technology for detection and phasing of structural variants in 13 ALL genomes. With this technology we were able to resolve nearly all of the aberrations detected by standard diagnostic analysis, and in the majority of the cases we were able to provide an improved delineation of the true structure of the chromosomal rearrangements. To achieve long haplotype blocks and thus optimal phasing of the diploid chromosomes, HMW DNA is recommended as source for library preparation. We found that, in our hands, standard column-based DNA extraction methods implemented in this project provided DNA molecules of sufficient length for accurate detection of structural variants that performed comparably to that of HMW DNA for detection of the known, subtype-defining translocations, albeit at the cost of shorter haplotype blocks. Consequently, this suggests that older biobanked DNA that has been extracted with standard columns may be used for genomic profiling by linked-read WGS. Although shorter haplotype blocks were obtained, we show that phasing information was not necessarily required to resolve several complex chromosomal rearrangements using standard DNA. Nonetheless, information about phased structural variants owing to longer haplotype blocks enabled detection of allele-specific alterations such as the abovementioned ERG and ETV6 deletions, thus enabling a more comprehensive delineation of the oncogenic lesions in ALL, consequently it would be highly recommended that HMW DNA is used when possible.

The rearrangement involving DUX4-IGH presents a particularly difficult aberration to resolve, and possibly why this fusion gene has not been identified until recently with RNA-sequencing studies. A likely explanation is the complex genomic region where DUX4 is located i.e. within the tandem repeat region D4Z4 consisting of 11-100 of repeat segments. The D4Z4 macrosatellite array is found within the subtelomeric region of chr4q and chr10q [141]. With the computational approaches used in this thesis, the insertion of DUX4 into the IGH locus was not be adequately resolved by linked-read WGS. Thus complementary assays, such as RNA-sequencing, were required to detect this aberration.

A particular challenge with linked-read WGS lies in the lack of tools that can accurately call structural variants with high confidence. In this thesis, we specifically focused on applying linked-read WGS to identify and characterize alterations that were already known to be present in the 13 ALL genome from diagnostic karyotyping or previous studies. In some instances we resolved previously undetected structural rearrangements; however they were
all related to a previously defined aberration. It is to be noted that many additional variants were called in each sample that we did not describe nor validate. Additional filtering or refined detection algorithms would be required to reduce the number of called structural variants in these data sets and to enrich for true somatic events. This may have been mitigated by including matched germline DNA in our study design, however this was not feasible for all of the patient samples included in the study. Also, all samples subjected to linked-read WGS had a high tumor purity (>90% leukemic blasts) and detection of aberrations in samples of lower tumor purity may be more challenging. Nonetheless, the linked-read WGS technology enabled simultaneous detection of aneuploidies, translocations and deletions in a single reaction, resulting in precise digital karyotypes, and thus holds great promise for future clinical incorporation.
Concluding remarks

In this thesis, by adopting a combined approach comprising DNA methylation classification, array-based copy number detection, fusion gene detection by RNA-sequencing and linked-read WGS, we could comprehensively dissect the genomic landscape of a large retrospective cohort of Nordic pediatric ALL patients. Collectively, these studies provide a more detailed view of the chromosomal abnormalities and expressed fusion genes in >1000 pediatric ALL patients than was possible based on the routine methods implemented at the time of ALL diagnosis. Many aberrations in ALL arise as a consequence of cryptic or complex rearrangements of the genome and are not detected at diagnosis by current diagnostic methods, such as karyotyping, SNP-arrays and FISH, which do not have the resolution necessary to comprehensively detect these complex rearrangements. Thus, new sequencing-based approaches are promising techniques to overcome this problem. Technical advancements in molecular profiling coupled with the reduction in sequencing costs are pivotal factors that make genome-wide sequencing more feasible for clinical diagnosis. Indeed, the simplicity of targeted panels for screening of a subset of specific aberrations may be enticing from a clinical context. However, owing to their limited genome-wide coverage, additional acquired alterations that are rare or even private, may not be fully resolved as compared to genome-wide approaches. Together, increased molecular understanding of the emerging subgroups is of great importance for further improved risk stratification and may be considered for future incorporation into clinical diagnostics of ALL.
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