

Molecular subtypes of gastrointestinal stromal tumors and their prognostic and therapeutic implications

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Gastrointestinal stromal tumors (GISTs) are composed of various molecular subtypes, with differing prognostic and predictive relevance. Previously, tumors lacking mutations in the *KIT* and *PDGFRA* genes have been designated as 'wild-type' GISTs; however, they represent a heterogeneous group currently undergoing further subclassification. Primary and secondary resistance to imatinib poses a significant clinical challenge, therefore ongoing research is trying to evaluate mechanisms to overcome resistance. Thorough understanding of the prognostic and predictive relevance of different genetic subtypes of GIST can guide clinical decision-making both in the adjuvant and the metastatic setting. Further work is required to identify tailored therapies for specific subgroups of GISTs wild-type for *KIT* and *PDGFRA* mutations and to identify predictive factors of resistance to currently approved systemic therapies.

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Gastrointestinal stromal tumors (GISTs) are the most common mesenchymal tumors of the digestive tract, representing 0.1–3% of all GI cancers. GISTs can arise from any part of the GI tract, primarily within the muscular wall of the stomach and small intestine and rarely in extra-intestinal locations (omentum, mesentery, retroperitoneum or pelvic cavity) [1]. The cells of origin of GISTs are thought to be the interstitial cells of Cajal or their precursors [2]. GISTs represent a wide spectrum of disease, with aggressiveness of the disease correlating with tumor size, mitotic activity and anatomical origin (these three clinicopathological features forming the basis of currently used risk-stratification systems) [1,3–5].

Surgery is the mainstay of management for localized GIST and is curative in 45–60% of cases [3,4]. Locally advanced or metastatic GISTs are notoriously refractory to conventional chemotherapy or radiation. The discovery of the *KIT* tyrosine kinase receptor and subsequently that of the mutually exclusive *KIT* and *PDGFRA* gain of function mutations have provided a paradigm shift in the way we classify, diagnose and most importantly treat GISTs. Studies of *KIT/PDGFRA* mutation negative or 'wild-type' (WT) GISTs have uncovered numerous other molecular groups, including mutations in *BRAF* and subunits of the succinate dehydrogenase (SDH) complex. Routine genotyping has become an integral part of management of GISTs undergoing tyrosine kinase inhibitor (TKI) therapy [5].

The objective of this manuscript is to mirror the evolution of GIST subclassification based on genetic profiling and to highlight the distinct prognostic and predictive relevance of already

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well-characterized and yet emerging genetic alterations in GIST. We separate in our discussion the prognostic and predictive relevance of specific genetic subtypes of GIST, therefore providing a more transparent guide for clinical decision-making in both the adjuvant and metastatic setting.

The KIT receptor tyrosine kinase

The *c-kit* proto-oncogene is allelic with the murine white-spotting locus (W), mutations of which affect melanogenesis, gametogenesis and hematopoiesis during development and in adult life [6]. *c-kit* encodes the 145-kDa receptor tyrosine kinase (RTK) KIT and is the normal cellular homolog of the viral oncogene *v-kit*, the transforming gene of the Hardy–Zuckermann 4 feline sarcoma virus [7]. KIT is a member of the type III RTK family which includes the PDGFRA and PDGFRB, the macrophage colony-stimulating-factor receptor (CSF1R) and the Fl cytokine receptor (FLT3) [8]. The CD117 antibody against KIT has previously been shown to be a sensitive and specific marker for GIST, being positive in 95% of GIST specimens [9].

The KIT transmembrane receptor is composed of an extracellular domain consisting of five immunoglobulin (Ig) like motifs, a transmembrane hinge, a juxta-membrane (JM) domain and an intracellular tyrosine kinase domain consisting of two regions separated by a kinase insert domain (KID). The different segments of the KIT receptor all have a specific designated role in the process of tyrosine kinase activation (Table 1).

In around 82–87% of all cases, GISTs have activating mutations in either the KIT or the homologous PDGFRA RTKs (Table 2). Gain of function mutation of either KIT or PDGRA receptors lead to constitutive, ligand-independent activation that results in the

activation of Ras/Raf/MAPK, JAK/STAT3 and PI3K/Akt/mTOR downstream pathways, ultimately increasing cell proliferation and inhibiting apoptosis [10,17].

• KIT exon 9, 11, 13, 17 mutant GIST

While most *KIT* mutations in GIST are somatic, germline mutations have been identified in a small number of families [5]. Gain-of-function mutations in *KIT* result in growth advantage by constitutive, ligand-independent activation of the RTK [18]. While most GISTs are heterozygous for a given mutation, in around 15% of tumors the remaining WT *KIT* allele is lost and this is associated with malignant behavior, increased mitotic activity and topoisomerase II expression [19].

Approximately 69–83% of all GISTs show a *KIT* mutation (Table 2). An important observation is that KIT detection by immunohistochemistry (IHC) is unrelated to the existence of underlying mutations. The vast majority of *KIT* ‘hot spot’ mutations are found in exon 11, less frequently in exon 9 and rarely in exon 13 and exon 17 (Table 2) [20]. Primary nonhot spot exon 8 mutation of the KIT receptor is extremely rare and is not routinely screened for (Table 2) [11].

• KIT exon 11 mutational landscape

The most common site of *KIT* mutation is in the 5′ end of exon 11, which encodes the JM domain, and in the overwhelming majority of cases deletions or substitutions involving codons 550–560 occur [12]. Genetic alterations in exon 11 disrupt the auto-inhibitory function and trigger ligand-independent receptor activation [21].

There is a constantly growing body of evidence that the exact types of genetic alterations hold strong clinical prognostic value of their own [13,14]. Wozniak and colleagues analyzed clinical follow-up data of 427 patients who

Table 1. KIT receptor function and primary mutational status in gastrointestinal stromal tumors.

Coding region	KIT segment	Physiological function	Primary mutation rate	Ref.
Exon 8	EC IGM	SCF/ligand binding; dimerization domain	0.15–0.23%	[10,11]
Exon 9	EC IGM	SCF/ligand binding triggering subsequent receptor homodimerization, conformational change and kinase activation	7–15%	[10,12–13]
Exon 11	JM	Auto-inhibition of the receptor in ligand-free state	61–71%	[10,12,14]
Exon 13	PKD	ATP-binding region	0.5–1.8%	[10,12,14]
Exon 14	KID	Undefined function beyond linking PKD with the DKD	NDA	[10]
Exon 17	DKD	Contains the activation loop that stabilizes the activated receptor	0.5–1%	[10,12,15]

NDA: No data available as primary mutation [10–16].
DKD: Distant kinase domain; EC IGM: Extracellular immunoglobulin-like motif; JM: Juxta-membrane domain; PKD: Proximal kinase domain; SCF: Stem cell factor.

Table 2. *KIT* and *PDGFRA* ‘hot spot’ mutation landscape.

Study (type)	OMR (%)	<i>KIT</i> (exon); %					<i>PDGFRA</i> (exon); %					Ref.
		All	9	11	13	17	All	18	18*	14	12	
Polish (registry)	82.2	69.3	7.3	61.1	0.5	0.5	12.9	11.9	8.2	0.7	0.2	[12]
ConticaGIST (registry)	85.1	71.1	7.4	71.1	1.8	0.6	14	12.8	9.8	0.3	0.9	[14]
EORTC 62005 (Phase III trial)	86.2	83.6	15.4	65.8	1.6	0.8	1.6	NS	1	NS	NS	[13]
CALGB 150105 (Phase III trial)	84.6	81.9	8.2	71.3	1.2	1.0	2.65	1.2	0.9	NS	0.23	[15]
ACOSOG Z9001 (Phase III trial)	87.4	76.2	6.9	67.3	1.8	0.2	11.2	NS	5.3	NS	NS	[16]

18*: *PDGFRA* exon 18 pD842V mutation; NS: Not specified; OMR: Overall mutation rate.

underwent curative resection for GIST in Poland between 1999 and 2009. Surgical specimens of imatinib-naïve GISTs from this period were prospectively included in the Polish Clinical GIST Registry. Mutational analysis was retrospectively carried out assessing for exon 9, 11, 13, 17 *KIT* and exon 12, 14, 18 *PDGFRA* mutational status [14]. The European ConticaGIST database study analyzed the clinicopathologic and molecular data (exon 9, 11, 13, 17 *KIT* and exon 12, 14, 18 *PDGFRA*) of 1,056 GIST patients undergoing curative R0/R1 resection. Patients were diagnosed between January 1985 and April 2012 with 13 contributing institutions from four European countries. As a strength of the ConticaGIST series, the majority of the cases (all those diagnosed from 2001) were studied prospectively (83% of all) [13]. None of the patients in either studies were exposed to chemotherapy or any other anticancer agent, including imatinib, thus both studies provide invaluable prognostic information on the clinical course of GIST according to specific tumor genotype.

Exon 11 deletions

Deletions affecting codons 557–558 of exon 11 of the *c-KIT* gene are detected in 23.2–27.7% of all GIST cases. They are lost either as specific isolated p.W557_K558 deletions in 6.3–7.5% of GISTs or as part of larger deletions in 15.7–21.4% of the cases (Table 3). Early studies associated deletions affecting codons 557–558 with an aggressive, metastasizing phenotype and indicate an overall poor prognosis [22,23]. Interestingly Martin-Broto and colleagues demonstrated that the predictive value of deletion of 557/558 for recurrence might be limited only to the first 4 years after curative surgery [24]. In the Polish registry study 557/558 codon deletions were more frequent in larger (88%, >5 cm) GISTs with higher mitotic index (MI; 75% with >5/50 HPF) and thus 80% of them stratified as high-risk tumors. Patients with 557/558 codon deletions had a lower 23.8% 5-year relapse-free survival (RFS) rate as

compared with patients with any other *KIT* exon 11 mutations (41.8% RFS), but also with other exon 11 deletions that have not involved codons 557/558 (33.3% RFS) [14].

In the Multinational European ConticaGIST Database analysis *KIT* p.W557_K558del mutants equally segregated in gastric and nongastric sites (55 vs 45%). *KIT* p.W557_K558del was more frequently identified in patients younger than 60 years of age (59 vs 42.4%), in tumors >5 cm (84.5 vs 57.7%), with MI >5/50 HPF (68.9 vs 39.4%), and classified as high risk (70.2 vs 38.9%), when compared with other *KIT* exon 11 mutated tumors. It was an important observation that the clinicopathologic characteristics of tumors bearing *KIT* p.W557_K558del were comparable with the group of tumors with *KIT* delinc557/558, within which tumor size, mitotic rate and fraction of high-risk tumors were also significantly higher than in tumors with other *KIT* exon 11 mutants. *KIT* delinc557/558 was associated with an increased risk for tumor progression with a hazard ratio (HR) of 1.45 and an inferior disease-free survival (DFS; median DFS 45.5 months; 5-year DFS 33.1%). The relatively high number of *KIT* del-inc557/558 mutants equally distributed in gastric and nongastric sites enabled the researchers to analyze the possible impact of this genotype on DFS, depending on the anatomical site of GISTs. In clear contrast with other *KIT* exon 11, *KIT* exon 9 and *PDGFRA* exon 18 mutations, the poor prognostic impact of *KIT* del-inc557/558 on patients' survival was only significant in GIST localized to the stomach ($p < 0.001$), but not in tumors with nongastric origin ($p < 0.26$). The same associations were also evident when comparing *KIT* del-inc557/558 mutants with other exon11 *KIT* mutations, overall ($p < 0.0001$; or comparing gastric ($p < 0.0001$) and nongastric ($p < 0.599$) GIST. This phenomenon might be related to the observation that gastric GIST with *KIT* del-inc557/558 had a larger size (7 vs 5.8 cm) and a higher mitotic rate (6 vs 4/50 HPF) when

Table 3. Exon 11 mutations and their prognostic implications.

Genetic alteration	Study/mutation frequency	Clinical–pathological prognostic features	Ref.
p.W557_K558 deletion	Polish Registry 7.5% ConticaGIST 6.3% Norwegian Registry 5.3% EORTC 62005 6.9%	High-risk tumors Higher MI, larger (>5 cm) size Gastric: nongastric location 1:1 Younger age at presentation (<60 years) Lower RFS rate compared with all other (<i>KIT</i> exon 9 and <i>PDGFRA</i> exon 18) and to other exon 11 mutations Prognostic power seems to be confined to gastric location of GISTs	[12–15,25]
KITdelinc557/558	Polish Registry 15.7% ConticaGIST 21.4%	High-risk tumors Higher MI, larger (>5 cm) size Gastric: nongastric location 1:1 Younger age at presentation (<60 years) Lower RFS rate compared with all other (<i>KIT</i> exon 9 and <i>PDGFRA</i> exon 18) and to other exon 11 mutations Prognostic power seems to be confined to gastric location of GISTs	[12–15,25]
Intron 10/exon 11 junction deletions (resulting in p.K550_K558 deletion)	Polish Registry 1.4%	High-risk tumors Presumed aggressive/metastasizing clinical behavior	[14]
Single nucleotide substitutions	Polish Registry 15.5% ConticaGIST 19.7% CALGB 150105 1.6% EORTC 62005 1.8%	Lower MI, smaller (<5 cm) size Indolent clinical course Better 5-year RFS (as compared with <i>KIT</i> deletions or duplications)	[12–15]
Duplications	Polish Registry 7% ConticaGIST 6.6% EORTC 62005 1.5%	Exclusive gastric location Benign clinical outcome	[12–14]
Homo/hemizygous <i>KIT</i> exon 11 mutant	Polish Registry 4%	High risk of early metastatic disease Disseminated malignancy at presentation	[14]

GIST: Gastrointestinal stromal tumor; MI: Mitotic index; RFS: Relapse-free survival.

compared with other *KIT* exon 11 mutants. Consequently, the fraction of patients with gastric GIST harboring *KIT* del-inc557/558 who relapsed 5 years after surgery was twice as high as patients with other *KIT* exon 11 mutations (61 vs 29%). Even in tumors classified as nonhigh risk ([very]low and intermediate) and originating from the stomach, the presence of *KIT* del-inc557/558 remained an important prognosticator for poor outcome in comparison with other *KIT* exon 11 mutations, *KIT* exon 9 and *PDGFRA* exon 18 mutations [13].

Intron 10/exon 11 junction deletions

In the Polish registry series six GISTs (1.4% of all) had deletions affecting the intron 10/exon 11 junction, resulting in p.K550_K558 deletion as such. Among these handful GISTs all but one tumor was high risk or overtly malignant at presentation, suggesting a more aggressive clinical behavior of tumors with such deletion [14].

KIT exon 11 single nucleotide substitutions

A discrepancy in the frequency of *KIT* exon 11 substitutions between population-based studies

15.5–28.6% and tumor material from clinical trials 1.6–1.86% has been observed (Table 3). The under-representation of *KIT* exon 11 substituted GISTs in the advanced/metastatic GIST trials would suggest a more indolent clinical behavior. In a Norwegian population study, the presence of point mutations was associated with a low mitotic count [25]. In the Polish registry study tumors with *KIT* exon 11 substitutions were also characterized by low mitotic activity and average size <5 cm. Patients with *KIT* substitutions have also been shown to have a better 5-year RFS rate (50.7%) than those with *KIT* deletions or duplications (28.1 and 40.0%, respectively) [14].

Duplications in *KIT* exon 11

Duplications in *KIT* exon 11 were historically associated with gastric tumor location and female gender, and also linked with a favorable clinical course [26]. The results of the Polish registry study confirmed this association since 26 out of 30 GISTs with 3'-end internal tandem duplications were of gastric origin and 21 of them were found in women. A relatively

good clinical outcome has also been observed in patients with these tumors [14].

Homo/hemizygous *KIT* exon 11 mutation status

Loss of the WT *KIT* 11 allele and presence of homozygous *KIT* exon 11 mutations was strongly associated with a malignant clinical behavior in gastrointestinal stromal tumors [27]. In the Polish registry study 4% (17) of *KIT* exon 11 mutant GISTs lacked a WT allele [14], indicating a homo/hemizygous *KIT* status. Eleven of the 17 cases were associated with a high risk of metastatic disease or were already disseminated at the time of presentation, corroborating previous findings.

Exon 9 *KIT* A502_Y503 duplication

KIT mutations in exon 9 coding for the extracellular domain occur in 7–15% of GIST cases (Table 2). These mutations are believed to mimic the conformational change that the extracellular *KIT* receptor undergoes when ligand is bound. *KIT* exon 9 mutations characterized by A502-Y503 codon duplications are almost exclusively found in intestinal GISTs and for long they have been associated with a more aggressive phenotype [28].

In the Polish registry study *KIT* A502_Y503dup mutation was not only associated with a small intestinal origin (27 out of 31 tumors) and malignant behavior, but also with a male predominance (21 out of 31) [14].

In contrast to previous observations Künstlinger and colleagues in 2013, described for the first time that nearly 20% of *KIT* exon 9-mutated GISTs actually occur in the stomach or rectum. They provided evidence that exon 9-mutated GISTs metastasize significantly more often to the peritoneum than to the liver. Analyzing the data of over 1500 GISTs from their registry, *KIT* exon 9 mutations were neither associated with intermediate-risk/high-risk status nor overrepresented among metastatic lesions and thus they concluded that exon 9 mutations *per se* do not have a prognostic relevance [29].

In the ConticaGIST registry study when compared with *KIT* p.W557_K558del, *KIT* p.A502_Y503dup had a lower median mitotic rate, whereas there was no difference in tumor size for these mutants. Overall, both *KIT* exon 9 and *KIT* del-inc557/558 mutants were associated with relatively equal and inferior DFS (median DFS in both groups, 45.5 months; 5-year DFS,

37.9 and 33.1%, respectively). There is solid evidence that compared with gastric tumors, small intestinal tumors with similar size and mitotic activity have a markedly worse prognosis. Across the board GISTs localized outside of the stomach were larger, had a higher mitotic rate and had worse 5-year DFS (34.2 vs 58.1%) in comparison with gastric tumors. As mentioned above, comparison between tumors with *KIT* exon 9 and *KIT* exon 11 mutations (both, *KIT* del-inc557/558 and other *KIT* exon 11) of non-gastric origin did not show differences in tumor clinical behavior as assessed by survival analysis. Thus, the authors concluded that in extra-gastric sites, the worse prognosis of *KIT* exon 9 mutants is related to the tumor location itself rather than to an intrinsic aggressive biologic nature of this mutation. In further support for this hypothesis, six out of seven gastric *KIT* exon 9 mutants in the ConticaGIST series study were classified as nonhigh risk, with only one of these patients developing progressive disease (PD), following a relatively long DFS of 56 months [13].

Exon 13 & 17

It has been estimated that the frequency of exon 13 and 17 mutations is not higher than 1–2% [30]. In the Polish registry study *KIT* tyrosine kinase domain mutations involving exons 13 and 17 were found in two cases each, supporting the notion that these are uncommon mutations [14]. Mutations in exon 17 encoding the activation loop of the kinase seem to stabilize the active conformation. Primary mutations, such as that of K642E in exon 13 encoding the ATP-binding region are extremely rare and are speculated to interfere with the physiological auto-inhibitory function of the JM domain [30].

In a multicenter study of 54 cases with primary *KIT* exon 13 or exon 17 mutant GISTs, specifically among the *KIT* exon 17 mutants twice as many tumors arose from the small bowel than the stomach. Intestinal site of origin of tumors was also over-represented among the *KIT* exon 13 mutants as compared with population-based studies. Overwhelming majority of the *KIT* exon 13 or exon 17 mutants displayed pure spindle-cell morphology, very rarely associated with epithelioid features. Gastric *KIT* exon 13 mutant GISTs were slightly larger and of a higher risk group than gastric GISTs on average, whereas the behavior of small intestinal GISTs with *KIT* exon 13 or *KIT* exon 17 mutations did not differ from other small intestinal GISTs [30].

• *PDGFRA* mutant GIST

PDGFRA mutations are reported in 1.6–2.7% of GISTs in Phase III clinical trials enrolling patients with advanced disease and up to 12.9–14% of primary tumors in population studies (Table 2). The markedly lower representation of *PDGFRA*-mutated GISTs in clinical trial material can be easily explained by a comparatively benign clinical behavior of these tumors. Historically they were indeed correlated with a more indolent course of disease [31]. Moreover, *PDGFRA* mutant GISTs are almost exclusively (90–93%) of prognostically more favorable gastric origin. The most prevalent genotype is the p.D842V substitution involving the second kinase domain (which corresponds to exon 17 of *KIT*), detected in 60–65% of all *PDGFRA* mutated tumors (Table 2). *PDGFRA* and *KIT* mutations are mutually exclusive and activate similar downstream signal transduction pathways [32].

In the Polish registry study, most changes among *PDGFRA* mutated tumors were identified in exon 18 (11.9%), including the p.D842V substitution and with in-frame deletion or deletion/insertion of different lengths (9–15 bp) in 8.2 and 3.7% of the cases, respectively. Three cases (0.7%) showed substitution in exon 14 of *PDGFRA* (p.N659K), and one (0.2%) revealed deletion in exon 12 JM domain (p.S566_E571delinsR). Multivariate analysis in this study revealed that patients with tumors with mutations involving *PDGFRA*, and *KIT* exon 11 substitutions or duplications have lower risk of 5-year relapse when compared with patients with *KIT* deletions involving codons 557/558 [14].

In the ConticaGIST series among tumors with *PDGFRA* mutations, the most prevalent was the p.D842V substitution (9.8% of all mutations and 65.2% of the *PDGFRA* exon 18 mutations). *PDGFRA* exon 18 mutation status correlated with an extremely favorable disease outcome (median DFS not reached; 5-year DFS, 75%) in comparison with other mutations. *KIT* exon 9 and *KIT* delins557/558 mutations were associated with a significantly increased risk for tumor progression (HR: 1.47 and 1.45), in contrast with *PDGFRA* exon 18 mutation (HR: 0.23). There was no significant difference in DFS of *PDGFRA* p.D842V versus other *PDGFRA* exon 18 mutations. Notably, among gastric *PDGFRA* mutations, the vast majority that progressed (11 of 14) carried an exon 18 *PDGFRA* D842V substitution. While the

number of the tumors with *PDGFRA* exon 18 mutations originating from nontypical anatomical sites was too low for a conclusive analysis, the frequency of relapse was lower in gastric versus nongastric *PDGFRA* exon 18 mutated tumors (11.8 vs 25%), respectively [13].

It has been suggested that GISTs with the *PDGFRA* exon 14 mutation represent a subset of clinically favorable gastric tumors (exclusively gastric location) with almost exclusively epithelioid morphology [33].

• *KIT/PDGFRA* mutation-negative 'WT' GIST

Approximately 15% of adult GISTs do not have detectable mutations in *KIT* or *PDGFRA* and historically were simply referred to as 'WT' GISTs. *KIT/PDGFRA* WT GISTs express high levels of *KIT* and can arise from any part of the GI tract. Phosphorylated *KIT* is detectable in some of these tumors, suggesting *KIT* activation may still have a role in their pathophysiology [34].

• *SDH*-deficient 'WT' GIST

About half of all *KIT/PDGFRA* WT GISTs have inactivating mutations in the genes coding one of the four (*SDHA*, *SDHB*, *SDHC* and *SDHD*) subunits of the *SDH* complex. The *SDH* complex is located in the inner mitochondrial membrane and plays a role in the electron transport chain and the Szent-Györgyi–Krebs cycle by replacing succinate to fumarate. Either gene mutations in any member of the *SDH* complex or an as-yet-unknown mechanism destabilizes the *SDH* complex. *SDH* enzyme dysfunction leads to accumulation of succinate, resulting in HIF1- α stabilization and HIF1- α controlled oncogene transcription. In WT GISTs without *SDH* activity, upregulation of HIF1- α may lead to increased growth signaling through IGF1R and VEGFR [35].

Double hit inactivation of any components of the *SDH* complex destabilizes the entire complex, resulting in degradation of the *SDHB* subunit. Several international groups have demonstrated that all *SDH* mutations are reliably detected by *SDHB* loss on IHC, therefore, *SDHB*-IHC is currently the method of choice to test for *SDH* deficiency in *KIT/PDGFRA* WT GIST [35–37].

SDH gene germline mutations are features of the Carney–Stratakis syndrome, an inherited predisposition to multiple gastric GISTs and paragangliomas. Carney-triad is characterized by multiple gastric GISTs, paragangliomas and

pulmonary chondromas with no *SDH* gene germline mutations detected (despite a deficiency of SDHB immunoreactivity) [38]. However, most recently it was suggested that in rare occasions the Carney triad can be allelic to Carney–Stratakis syndrome. In the largest published cohort of 63 unrelated Carney triad patients six patients (9.5%) were found to have germline variants in the *SDHA*, *SDHB* or *SDHC* genes [39]. Detecting SDH deficiency therefore, will trigger further clinical investigations in order to exclude syndromic GISTs, especially in younger patients.

SDH-deficient GISTs are particularly common in childhood and young adulthood, approximately 1–2% of all GISTs occurring in the pediatric population [40]. These SDH-deficient pediatric GISTs are characterized by unique clinical, morphological and genetic features. They have a predilection for the stomach, commonly demonstrate a multilobulated/multinodular growth pattern, frequently metastasize to lymph nodes and are mostly WT for *KIT* and *PDGFRA* [35,36].

Importantly SDH-deficient ‘pediatric-type’ GISTs also account for between 5 and 7.5% of all gastric GISTs occurring in adults [35]. It must be emphasized that the prognosis of SDH-deficient GISTs cannot be predicted by size and mitotic rate as even small, mitotically inactive SDH-deficient GISTs may metastasize. Interestingly when metastases do occur they may be strikingly indolent, sometimes remaining stable for years or decades [37]. As their distinctive histology predicts genotype and clinical behavior it can be recommended that IHC for SDHB should be performed on gastric GISTs with compatible ‘pediatric-type’ morphology.

In a recently published study *SDHA* mutations were associated with statistically significant better clinical outcome as compared with *KIT/PDGFR*A mutations and *KIT/PDGFR*A WT without SDH deficiency. All survival analyses (from diagnosis of primary tumors and from diagnosis of metastatic disease) confirmed a far more indolent course of disease for patients with *SDHA* mutated WT GISTs [41].

SDH proficient ‘WT’ GIST

In the absence of the well-characterized frequent RTK mutations further, far less frequent mutations have been described in the members of the downstream signaling pathway, with mutations in *BRAF* (V600E), *HRAS*, *NRAS* or *PIK3CA* genes. These mutations presumably cause the

constitutive activation of KIT downstream signal pathways. In addition, *KIT/PDGFR*A WT GIST may be related to syndromic neurofibromatosis type I (NF-1) disease, associated with NF1 protein loss of function due to genomic inactivation of the *NF1* gene [5].

Hostein *et al.* screened 321 GISTs with 70 WT GISTs for *BRAF* mutation. Similar to other tumor types where *BRAF* mutations are more commonly observed, the mutations seen in GIST are also located within the exon 15 V600E hot spot. *BRAF* V600E was detected in nine (13%) of the 70 WT GISTs and no mutations were detected in GISTs bearing *KIT* or *PDGFRA* mutations. Interestingly *BRAF* V600E detection in the tumor did not result in a higher expression of the B-raf protein or the preferential activation of the p42/44 MAPK signaling pathway compared with GISTs without the *BRAF* mutation [42]. *KIT*, *PDGFRA* and *BRAF* WT GISTs are commonly referred to as ‘triple-negative’ GISTs.

Miettinen and colleagues noted that patients with NF1 are overrepresented by at least 45-fold among GIST patients and GISTs occur in approximately 5–25% of NF1 patients. NF1-associated GISTs develop secondary to a somatic inactivation of the WT *NF1* allele in the tumor and are commonly multicentric, predominantly located in the small intestine and lack *KIT* and *PDGFRA* mutations. The majority of NF1-associated GISTs present as small, low mitotic index lesions and they are associated with quite favorable long-term clinical outcomes reflected in low recurrence and metastases rates. Interestingly NF-1 associated GISTs arising from the duodenum display an aggressive behavior, being large mitotically active tumors with pronounced metastatic potential [43].

A small subgroup of *KIT/PDGFR*A WT GIST, referred as ‘quadruple WT GIST’ (Q-WT GIST), that lack mutations in any of the known *KIT* exons (8, 9, 11, 13, 14, 17) or *PDGFRA* exons (12, 14, 18) or RAS pathways, including *BRAF* (exons 11, 15) and *RAS* (exons 2, 3), or *NF1*, and yet retain an intact SDH complex (SDHB IHC positive, and no mutations in *SDH*) has been identified. The true clinical relevance of this subgroup is yet to be elucidated [44].

Most recently two oncogenic RTK translocations were reported in a subset of nongastric Q-WT GISTs [45,46]. Two cases with ETV6-NTRK3 fusion (in a colonic and a rectal Q-WT GIST) and one with FGFR1-TACC1

translocation (in a small intestinal Q-WT GIST) were found. Both of these alterations have been identified in other malignancies and are known to constitutively activate the target kinases. These RTK translocations found in a subset of Q-WT GIST seem to define a novel biological class of GIST arising in nongastric sites. The oncogenic RTK fusion proteins could represent potential therapeutic targets. If confirmed in larger series, routine testing for RTK translocation may be indicated for Q-WT GIST [45,46].

• Genetically unstable GIST

The accumulation of chromosomal abnormalities correlates with the biological behavior of GISTs. About two-thirds of GISTs exhibit 14 monosomy or partial loss of 14q, and half of them have loss of the long arm of chromosome 22. Chromosome 14 or 22 aberrations are linked to a borderline malignant potential. An aggressive biology is associated with the loss of chromosome 1p, 9p (spanning CDKN2A or p16INK4A) and 11p regions. Gain in chromosomal segments 8q and 17q have been associated with increased metastatic potential. It is an interesting observation that while an unstable karyotype correlates with the presence of GIST mutations, *KIT/PDGFR* WT GISTs are genomically stable [47,48].

Combined prognostic value of mutational status in treatment-naïve GIST

As detailed above individual genetic alterations, for instance *KIT* exon 11 deletions, were reported already a decade ago as an independent adverse prognostic factor in patients with untreated GIST [11]. However, until very recently, the combined prognostic value of the most relevant genetic GIST subtypes has not been elucidated. Rossi *et al.* analyzed a series of 451 untreated primary localized GISTs for *KIT*, *PDGFRA* and *BRAF* mutations [49] and found that mutational status is a significant prognostic indicator of overall survival (OS) in treatment-naïve, localized GISTs. Patients with *KIT*-mutated tumors had a worse outcome than *PDGFRA*-mutated or ‘triple-negative’ (*KIT*, *PDGFRA*, *BRAF* WT) cases. Based on multivariable Cox regression models the authors identified three distinct molecular risk groups. Group I, consisting of *PDGFRA* exon 12, *BRAF* and *KIT* exon 13-mutated cases, exhibited the best clinical outcome. The intermediate risk (HR: 3.06) Group II, included ‘triple-negative’,

KIT exon 17, *PDGFRA* exon 18 D842V and *PDGFRA* exon 14-mutated GISTs. Group III, comprised of *KIT* exon 9 and exon 11 and *PDGFRA* non-D842V exon 18 mutant GISTs, displayed the worst clinical outcome (HR: 4.52). This study clearly highlighted the prognostic impact of mutational status on the natural history of GIST. Inclusion of molecular prognostic grouping into currently used clinicopathologic risk stratification criteria could clearly fine tune the decision-making process for adjuvant therapy.

• Therapeutic implications of different genetic subtypes of GIST

Due to its ubiquitous role in the pathogenesis of GISTs, *KIT* has become a universal therapeutic target. Imatinib, a competitive inhibitor of the ATP-binding domain can only bind to the inactive conformation of *KIT* and its pronounced clinical efficacy was confirmed one and a half decades ago [50,51]. Imatinib represents the standard upfront medication for the treatment of advanced/metastatic GIST, with an overall 80% disease control rate (objective response or stable disease), a median progression-free survival (PFS) of approximately 20 months and a median OS of around 50 months [52,53].

Over the last decade *KIT* mutational status has emerged as a strong predictive indicator of treatment response. GISTs with *KIT* exon 11 mutated genotype show a marked, approximately 70% objective response rate (ORR; complete response/partial response) as compared with the 40–45% ORR seen in exon 9 mutant and WT GISTs. The ORR advantage seen in exon 11 mutant patients translates into an additional OS gain of approximately 20 months for this subgroup compared with exon 9 mutated and ‘WT’ GISTs (60 vs 40 months for exon 9 mutant and WT GIST, respectively) [52,53].

In the pooled analysis of the two pivotal Phase III trials comparing 400 versus 800 mg daily imatinib dose, the sole predictive factor of response was the presence of a *KIT* exon 9 mutation. The estimated risk of progression or death was reduced by 42% in the high-dose arm (compared with the standard-dose arm) in patients with *KIT* exon 9 mutated tumors. However, no significant difference in OS was seen between patients treated with 400 and 800 mg imatinib, irrespective of mutational status. The study concluded that for most patients, the recommended daily dose is 400 mg daily, with the exception of

KIT exon 9 mutated tumors where the 800 mg dose can be considered [54].

Approximately, 5% of patients with GIST do not express *KIT*, but may have tumors that harbor imatinib-sensitive *KIT* or *PDGFRA* mutations; therefore, patients with *KIT*-negative GISTs should not, *a priori*, be denied imatinib therapy and tumor samples should be sent for mutational analysis [55].

Despite majority of GIST patients benefiting from imatinib treatment, approximately 10–15% of them show primary resistance with early progression within 3–6 months of initiating therapy, in keeping with primary resistance. Although 80–85% patients with advanced GIST benefit from imatinib treatment, 40–50% of patients subsequently develop secondary resistance to the agent with a median time to progression of about 24 months [20,52,56].

• Primary imatinib resistance

The accumulated results of preclinical and clinical research have provided powerful tools in the explanation, prediction and management of primary resistance. Both primary and secondary resistance to imatinib can be partially explained by a conformational shift in the kinase domain of *KIT* and *PDGFRA* that favors the activated state [51]. Preclinical data indicated that *PDGFRA* isoforms with a substitution involving codon D842 in exon 18 (D842V, RD841–842KI, DI842–843IM) lead to primary resistance to imatinib, with the exception of D842Y. Other mutations in exon 18 (D846Y, N848K, Y849K and HDSN845–848P) were found to be imatinib sensitive. Imatinib can only bind to the inactive conformation of both the *KIT* and *PDGFRA* receptors. *PDGFRA* D842V mutation results in a distortion of the kinase activation loop, thus strongly tilting the protein conformation in favor of the activated structure. In clinical trials *PDGFRA* mutant GISTs showed a mild sensitivity to drug (66%) except the exon 18 D842 V mutation which proved to be resistant [15,57]. However, a most recent large multicenter observational study reported objective response to imatinib in a small proportion of patients with *PDGFRA* D842V-mutated GIST. Out of 16 patients with the mutation 12.5% had partial response, 18.8% had stable disease and 56.3% had PD as best response according to Choi criteria. Median time to progression was 8.0 months (range: 0–42). The authors of the report rightly concluded that as patients with

PDGFRA D842V-mutated GIST do have a small chance of responding to standard TKI treatment, imatinib should not be universally denied in patients harboring this mutation [58].

It is currently unclear how the predictive power of genetic alterations seen in the advanced setting may translate into decision-making regarding adjuvant therapy. The two adjuvant trials (the USA ACOSOG Z 9001 and the German–Scandinavian study) have failed to demonstrate that a specific mutation can predict a better RFS and OS in treated patients [16,59].

In the ACOSOG Z 9001 trial on multivariable analysis of patients tumor genotype was not significantly associated with RFS, however patients with *KIT* exon 11 deleted tumors assigned to 1 year of adjuvant imatinib did have a longer RFS. The excellent survival of patients with tumors harboring *PDGFRA* mutations in the placebo group provides a good argument that these patients may not require adjuvant treatment [16]. In the German–Scandinavian study *KIT* exon 11 mutant GIST patients benefited from the longer 3-year treatment, whereas no significant improvement over 12 months of imatinib was found in *KIT* exon 9 or *PDGFRA* mutant or WT patients (note that the number of patients were small in these later categories) [59].

In a small series of seven metastatic NF1-associated GIST patients, three of the four imatinib-treated patients showed primary resistance to the treatment (all three tumors were ‘WT’ GISTs). The fourth metastatic patient with an exon 18 mutated tumor had temporary stable disease. Median OS for this four-patient cohort was 21 months [60].

Rege and colleagues in their seven patient cohort of metastatic ‘pediatric-type’ SDH-deficient adult GISTs reported absolute primary resistance to imatinib in all of the cases. In contrast to this, in a more recent 2015 report 4/5 of imatinib-treated ‘pediatric-type’ SDHA-mutant metastatic GISTs showed a longer than 6-month PFS, with two ongoing responses at 19 and 58 months [61]. This later observation begs the question whether SDHA mutational status shall be performed routinely in SDH-deficient tumors as assessed by IHC.

Alternate signaling pathway mutations (like *BRAF* exon 15 activating mutation) in patients lacking identifiable *PDGFRA* or *KIT* mutations can be potential alternative mechanisms explaining their frequent primary resistance to imatinib [62–64].

- **Secondary imatinib resistance**

Acquisition of *KIT* or *PDGFRA* secondary mutations represent the most frequent mechanism of imatinib resistance in GIST [64,65]. Radiological evidence of clonal resistance can be detected as the appearance of one or more areas of increased vascularity within a previously responding or stable lesion. These lesions can precede by several months (PD) according to Response Evaluation Criteria In Solid Tumors (RECIST). In a study of TKI pretreated patients [66], the mutational status of individual nodules within a dominant tumor mass confirmed intra- and inter-lesional mutational heterogeneity between co-existing nodules. In total, 83% of samples had at least one secondary mutation, 67% had two to five different secondary mutations in different tumor samples and 34% had more than two mutations in the same clone. Secondary mutations clustered in the *KIT* ATP-binding pocket (exon 13) and kinase activation loop (exon 17). *KIT* amplification was detected by FISH in two of the ten metastases lacking secondary *KIT* mutations. No *KIT* kinase resistance mutations were detected in *KIT/PDGFR* WT GISTs or in *KIT*-mutant GISTs showing unusual morphology and/or loss of *KIT* expression by IHC. These results clearly expose the emergence of heterogeneous resistance mechanisms in these tumors. Furthermore, these data indicate that repeat biopsy (with mutational analysis) may not be relevant in metastatic GIST and highlight the therapeutic challenges involved in managing heavily pretreated patients [66]. Emergence of polyclonal imatinib resistance poses a great difficulty in developing effective single agent next-generation kinase inhibitors.

Secondary *KIT* exon 17 mutations affecting the activation loop stabilize the active receptor conformation, while imatinib can only bind and inhibit the nonactivated (auto-inhibited) conformation of *KIT*. Activation loop mutations therefore indirectly induce imatinib resistance by shifting the equilibrium strongly in favor of the active conformation [21].

In a Phase I/II trial half of the patients progressing on imatinib had a secondary *KIT* mutation [67]. Mutational distribution was nonrandom, secondary mutations clustering in exons 13 and 14, encoding the drug/ATP-binding pocket of the receptor and in exon 17, encoding the kinase activation loop. The most commonly detected secondary mutation was the exon 13 V654A. Two tumors had secondary mutations in

KIT exon 18. One patient had distinct secondary mutations in separate lesions, that of exon 13 V654A and exon 17 D816H respectively. Of the four primary *PDGFRA* mutant samples one had a secondary mutation in exon 18 (primary mutation in exon 12), two lacked secondary mutations (both had primary exon 18 D842V mutations) and in the fourth case there was no postimatinib sample available. More interestingly, in the post-imatinib samples of eight *KIT* or *PDGFRA* wild type GISTs no secondary mutations were found. Secondary mutations were more likely to be found in patients who initially harbored *KIT* exon 11 mutations (73%) as compared with *KIT* exon 9 mutations (19%). The higher rate of secondary mutations in the primary exon 11 mutant subgroup, is likely to be due to the longer exposure to imatinib. In preclinical *in vitro* studies sunitinib potently inhibited the kinase activity of *KIT* receptors that contained imatinib-resistant secondary mutations in the drug/ATP-binding pocket, such as V654A (exon 13) and T670I (exon 14). Furthermore, *in vitro* sunitinib was relatively ineffective at inhibiting *KIT* receptors with secondary mutations affecting the activation loop. In keeping with these findings, clinical trial data have shown that PFS and OS were longer and the clinical benefit rate was higher for patients with tumors harboring the *KIT* exon 13 or 14 ATP-binding-pocket mutations than those with *KIT* exon 17 or 18 activation loop mutations [67].

Amplification of the *KIT* and/or *PDGFRA* genes has also been postulated as a potential mechanism for either primary or delayed TKI resistance [67,68]. Some GISTs lacking secondary kinase mutations do show genomic amplification of *KIT* and/or become hemi- and/or homo-zygous for the primary *KIT* mutation undergoing deletion of the WT *KIT* allele [68].

IGF1R amplification may represent another mechanism of *de novo* or acquired imatinib resistance. It has to be noted that IGF1R overexpression was detected in the overwhelming majority (89%) of SDH-negative gastric GISTs but in only 1% of SDHB-positive gastric GISTs [69].

- **Treatment beyond imatinib**

Sunitinib and regorafenib have been approved for imatinib-resistant GIST based on placebo controlled randomized Phase III trials [70,71].

The response to second-line sunitinib correlates with the primary (pre-imatinib) tumor

mutation status [67,72]. In a worldwide, open-label treatment-use study of 1124 patients resistant or intolerant to imatinib a median PFS of 7.1 months was reported on sunitinib. The PFS in patients with a primary *KIT* exon 9 mutation was significantly better as compared with those with primary exon 11 mutation, with figures of 12.3 and 7.0 months, respectively. Moreover, primary *KIT* exon 9 mutation was associated with a longer OS and higher ORR as compared with exon 11 GISTs [72].

As far as secondary *KIT* mutations are concerned, for primarily *KIT* mutant GIST patients the median PFS and OS with sunitinib was significantly longer for secondary *KIT* exon 13 or 14 mutations than for those with secondary exon 17 or 18 mutations [67]. These results corroborate the *in vitro* findings which showed that sunitinib effectively inhibits the phosphorylation of *KIT* double mutants where secondary mutation affected the drug/ATP-binding pocket. On the other hand, sunitinib has very little activity against *KIT* double mutants where secondary mutations affected the activation loop [73].

Moreover, preclinical studies of imatinib-resistant GIST cell lines evaluating sunitinib sensitivity showed that only *KIT* T670I (gatekeeper) and V654A mutants were sensitive to sunitinib, while *PDGFRA* D842V (activation loop) mutants remained resistant [73]. Unfortunately, the available clinical data are yet too limited to investigate the effects of *PDGFRA* mutations on efficacy outcomes following sunitinib treatment [67,72].

There is little evidence available in regard of the predictive power of mutational status in the third-line regorafenib treatment of GISTs; however, regorafenib showed efficacy in all genetic subtypes of GISTs including exon 9 and *PDGFRA* D842V mutants [71,74].

Conclusion

The discovery of activating mutations in *KIT* and *PDGFRA* and their role in the pathogenesis of GIST has revolutionized our understanding of the biology and therapy of this disease. The introduction of imatinib has ultimately led to a radical improvement in life expectancy of advanced GIST patients. However, it is clear that GISTs are composed of many different molecular subtypes, with differing clinical characteristics and response to therapy. The detection of specific somatic genetic changes in GISTs can provide potential predictive and prognostic

information for guiding therapy. We can conclude that the term ‘WT’ GIST is outdated by a series of recent findings, and in case it is used reference always should be made which genes it applies to (i.e., *KIT/PDGFR*A WT GIST or *KIT/PDGFR*A/*BRAF* WT GIST and so on).

Nonhot spot exon 8 and exon 14 mutations of the *KIT* receptor are extremely rare and are not routinely screened for. There is a genuine concern that without identifying these rare mutations we miss to identify vital information about the biological behavior and treatment sensitivity (both in the adjuvant and metastatic setting) of these genetic subtypes. *KIT* exon 11 mutant patients are twice as likely to respond to imatinib than those with exon 9 mutant or WT GISTs. Moreover, higher response rates to imatinib in the *KIT* exon 11 mutant group translates into a PFS and OS advantage as compared with all non-*KIT* exon 11 mutant GISTs. The presence of a *KIT* del-inc557/558 mutation in a study of patients from the pre-imatinib era, reported that this mutation was strongly associated with poorer outcomes both in high-risk and non-high-risk gastric GIST, indicating its additional prognostic value for patient selection for adjuvant therapy. It remains to be shown whether patients with GISTs with *KIT* del-inc557/558 genotype should not be kept longer (or even life-long) on standard adjuvant imatinib treatment (i.e., 3 years).

KIT exon 9 mutations have been associated with a poor clinical prognosis as compared with other mutations. However, the worse prognosis of *KIT* exon 9 mutants seems to be related to the almost exclusive high-risk extra-gastric tumor location of these GISTs rather than to an intrinsic aggressive biologic behavior. *KIT* exon 9 mutant GISTs are more likely to respond to the higher 800 mg dose of imatinib, and in the advanced setting patients there is a strong recommendation of starting patients with tumors harboring these mutations on the higher dose. The optimal dose of imatinib in the adjuvant setting for *KIT* exon 9 mutant GISTs is yet to be elucidated. Identification of *PDGFRA* mutations in GIST proved to be a strong predictor of good clinical outcome and this molecular factor could therefore add a significant value to the current consensus risk criteria used for GIST stratification. In addition, given that these mutations comprise the majority of cases with the imatinib-resistant p.D842V subtype, mutational testing is highly relevant in order to avoid overtreatment

of gastric tumors with imatinib in the adjuvant setting. As tumors with certain mutations such as *PDGFRA* D842V mostly have primary resistance to imatinib and sunitinib, patients should be offered participation in clinical trials of novel agents. A recent large multicenter observational study has suggested that some tumors with the D842V mutation may respond to standard tyrosine kinase therapy. While tumors with no *PDGFRA* or *KIT* mutations (WT GISTs) are generally considered to be less responsive to these agents, more focus should be directed at the exact mutational landscape of these cancers with regard to the therapeutic decision-making process. These are important observations that should be factored into the decision-making process prior to commencing tyrosine kinase therapy.

Future perspective

Inevitably all currently available standard treatment options lose their efficacy to control advanced GIST. Ongoing preclinical and clinical research is focusing on evaluating novel

therapeutic approaches to overcome primary and secondary resistance to imatinib and the other two currently available licensed medications, sunitinib and regorafenib. Targeting deregulated downstream pathways shall provide further treatment options in the management of imatinib/sunitinib/regorafenib insensitive/resistant GISTs.

Emerging information about the prognostic value of different, specific mutations warrants further evaluation, possibly using pooled cohorts stratified for the mutational status from the available prospective clinical trials in the adjuvant setting.

Currently, adjuvant therapy in ‘high-risk’ WT GIST and the optimal systemic treatment for metastatic WT GIST remain debatable ‘hot topic’ questions, with no clear-cut guidelines. The group of WT patients is heterogeneous and further attempts should be made to further delineate individual molecular subtypes, in order to guide future clinical trial development and optimize systemic therapy.

EXECUTIVE SUMMARY

Classification of gastrointestinal stromal tumors

- Gastrointestinal stromal tumors (GISTs) are the most common type of mesenchymal tumors of the digestive tract, mainly defined by the presence or lack of mutually exclusive gain-of-function mutations in the *KIT* and *PDGFRA* receptors.
- GISTs historically referred to as ‘wild-type’ (WT) GISTs are a very heterogeneous group which do not have a detectable mutation in either the *KIT* or the *PDGFRA* receptor genes, while *KIT* activation by phosphorylation is still detectable in these tumors.
- WT GISTs can be split into two large groups, based on whether they are proficient or deficient in the succinate dehydrogenase complex.
- Detecting mutations in alternative pathways can further subclassify WT GISTs.

Therapeutic implications of genetic subtypes

- *KIT* exon 11 mutant patients are twice as likely to respond to imatinib than those with exon 9 mutant or WT GISTs with a progression-free and overall survival advantage as compared with all non-*KIT* exon 11 mutant GISTs.
- *KIT* exon 9 mutant GISTs are more likely to respond to the higher 800 mg dose of imatinib, and in the advanced setting patients can be started *ab ovo* on the higher dose.
- The *PDGFRA* D842V isoform with a substitution involving codon D842 in exon 18 is generally believed to lead to primary imatinib resistance. Caution should be exercised when it comes to therapeutic decisions as recent data suggest some response to imatinib in this subset of tumors.
- While *SDH*-deficient ‘pediatric-type’ GISTs have been previously attributed absolute primary imatinib resistance, most recent reports suggest imatinib responsiveness in *SDHA*-mutated tumors.
- Acquisition of secondary mutations in either *KIT* or *PDGFRA* represents the most frequent mechanism of imatinib resistance in GIST.
- Adjuvant therapy in ‘high-risk’ WT GIST and the optimal systemic treatment for metastatic WT GIST remain debatable ‘hot topic’ questions, with no clear-cut clinical guidelines.

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