

Chronic Graft-Versus-Host Disease

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44.1 Introduction

Chronic GVHD (cGVHD) is the most relevant cause of late non-relapse morbidity and subsequent mortality (approximately 25%) following allo-HSCT (Grube et al. 2016). Its incidence is approximately 50% among all patients following allo-HSCT and has increased during the last two decades due to increasing patient age and increasing use of unrelated and/or mismatched donors, RIC regimens, and PBSC (Arai et al. 2015). While the incidence of cGVHD is lower (20–40%) in children, its incidence rises to 60% as age increases (Baird et al. 2010).

The pathophysiology of cGVHD is different from aGVHD and mainly characterized by impaired immune tolerance mechanisms affecting innate and adaptive immunity. Both autoreactive and alloreactive donor-derived T and B cells play a role (Cooke et al. 2017). Other pathophysiological factors are indirect presentations of alloantigens through antigen-presenting donor cells and mechanisms of chronic inflammation with subsequent scar formation and fibrosis. One important aspect of GVHD pathophysiology is the variability of immune reconstitution, which is

body irradiation (Baird et al. 2010). By far the strongest predictor is the history and severity of acute GVHD.

In addition to the harm it causes, cGVHD also has a *protective effect*, as patients with cGVHD have lower rates of recurrence of their underlying malignant disease (Grube et al. 2016). Overall survival of patients transplanted for malignant diseases developing mild cGVHD

nonmalignant diseases.

also has a *protective effect*, as patients with cGVHD have lower rates of recurrence of their underlying malignant disease (Grube et al. 2016). Overall survival of patients transplanted for malignant diseases developing mild cGVHD is therefore better compared to patients without cGVHD. Even OS of patients with moderate cGVHD is not different from patients without cGVHD, as the slightly increased mortality associated with cGVHD is counterbalanced by lower disease-associated mortality (Kuzmina et al. 2012).

age-related and dependent on thymic function and hormones. This adds to the unpredictability

of the effects of transplant procedures and com-

plications in a very heterogenous cohort of chil-

dren and adolescents with malignant and

cGVHD are unrelated and/or mismatched donor.

PBSCs as donor source, older donor age, female

donor into male recipient, and the use of total

Known risk factors for adult and pediatric

In contrast, the *long-term mortality* rate of patients with severe cGVHD is as high as 50% taken into account that the severity is less relevant compared to certain risk factors for mortality consisting of low platelets at diagnosis of cGVHD, the direct progression of acute GVHD into cGVHD (progressive onset), and certain organ manifestations

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(lung, gastrointestinal and cholestatic liver involvement) (Grube et al. 2016). One important pediatric aspect involves the high proportion (up to 50%) of nonmalignant underlying diseases as HSCT indication. While malignant diseases benefit from the graft-versus-malignancy effect induced by GVHD, it only offers harm for the nonmalignant diseases. In daily clinical routine, this fact influences GVHD prophylaxis and treatment both in regard to intensity and duration of immunosuppressants (Lawitschka et al., data of a survey by the EBMT pediatric diseases WP, submitted). However, prospective pediatric data of immune reconstitution in GVHD patients evaluating the influence of underlying diseases are scarce.

44.2 Clinical Manifestations

cGVHD usually begins between 3 months and 2 years after HSCT, but earlier onset (at least 1 month after transplantation) is possible (Jagasia et al. 2015). Besides classical manifestations, cGVHD can imitate almost any autoimmune disease, such as myasthenia gravis and myositis. As cGVHD can affect a number of organs, and patients often do not report changes until functional impairment is recognized, regular examination of all organs potentially affected is essential. The following section describes the most common clinical organ manifestations of cGVHD. In general, pediatric manifestations are similar to adult cGVHD; when indicated, specific aspects are shortly described.

44.2.1 Skin

The skin is the most frequently involved organ with different morphology, depending on the different skin layers (epidermis, cutis, subcutis, and fasciae) involved. Some manifestations may overlap with acute GVHD like erythema, maculopapular rash, and pruritus. Cutaneous cGVHD may show many different non-sclerotic and sclerotic phenotypes often simulating well-known chronic inflammatory and autoimmune diseases (Strong Rodrigues et al. 2018).

Diagnostic features of NIH-defined cGVHD include poikiloderma, lichen planus-like, lichen

sclerosus-like, morphea-like, and deep sclerotic eruptions, and no biopsy is needed to confirm the diagnosis. Distinctive for cGVHD, other or common skin manifestations like depigmentation and papulosquamous lesions or ichthyosis, keratosis pilaris, pigmental changes, loss of skin appendages, and sweat impairment are not sufficient for diagnosis and require histopathological confirmation if no diagnostic signs in the skin or other organs are present (Jagasia et al. 2015).

In pediatric patients, the incidence of viral reactivation and infection seems higher (although only proven for some viruses), and therefore infection has to be ruled out. Viral skin infections can worsen or activate cGVHD (Jacobsohn 2010). Premature graying of the hair is even in small children common, possibly together with seborrheic scalp changes. Of note, if sweat glands are destroyed, this may be of importance for phototherapy because of the inability to sweat with consequent hyperthermia.

44.2.2 Eyes

cGVHD of the eyes usually manifests as keratitis sicca. In addition to atrophy of the lacrimal gland with subsequent tear deficiency (sicca syndrome), the meibomian glands and eyelids are often affected by severe blepharitis which may initially present with tearing. Around the conjunctiva there are often not only fibrotic alterations but also chronic persistent inflammation with visible erythema of the conjunctiva. As dry eye symptoms are rarely communicated by children, light sensitivity is the predominant symptom, sometimes with excessive eye rubbing. Infections have to be ruled out. Referral to a pediatric experienced ophthalmologist is recommended.

44.2.3 Oral Mucosa

Oral manifestations may appear as erythema or lichenoid changes (the latter are regarded as diagnostic) of the oral mucosa as well as ulcera and mucoceles. Sicca symptoms may result from destruction of the salivary glands. Long-term cGVHD may lead to gingivitis, periodontitis, increased tooth decay, and tooth loss. In children

excessive drinking during eating may be the first symptom of oral involvement. Not only mucosal problems but abnormal teeth development (e.g., hypodontia, root malformation, enamel hypoplasia) and caries are often seen as secondary symptoms in infants.

44.2.4 Liver

Liver involvement manifests as cholestasis and may resemble primary biliary cirrhosis, but hepatitic forms with high transaminases are also possible. Other factors, such as viral infections (hepatitis A, B, C, and E, CMV, EBV, ADV, and HHV6/7), drug toxicity, or total-parenteral nutrition-related cholestasis, should be excluded, but liver biopsy may be required to confirm the diagnosis, particularly in patients with no other symptoms of cGVHD and failure to respond to initial treatment of suspected GVHD (Stift et al. 2014).

44.2.5 Gastrointestinal Tract

GI manifestations can lead to dysphagia (esophagus), nausea and vomiting (stomach), or chronic diarrhea and malabsorption syndrome (intestines, pancreas). Occasionally cGVHD may also manifest as immune-mediated pancreatitis. Of note, except esophageal involvement, intestinal involvement is regarded as manifestation of acute GVHD, and patients are therefore classified as suffering from overlap syndrome in which concomitant symptoms of chronic and acute GVHD occur.

Infections like ADV or CMV gastroenteritis, secondary gluten or lactose intolerance, pancreatic insufficiency, and drug-related side effects (e.g., mycophenolate mofetil) have to be ruled out.

Malnutition and enteral fluid and protein loss in small children require regular laboratory monitoring.

44.2.6 Genitals

The symptoms of cGVHD are similar to those of genital lichen planus which may occur in males and females. Vaginal synechiae, ulceration, and fissures can subsequently occur. Genital manifestations are often associated with oral manifestations of cGVHD. As symptoms may not be reported spontaneously, females suffering from cGVHD require regular gynecological follow-up. In girls cGVHD may manifest with vulvovaginitis, in boys with balanitis or balanoposthitis. Of note, healing may occur with fibrosis possibly leading to synechia with the risk of hematocolpos during puberty in females and of phimosis in males.

44.2.7 Lung

Pulmonary manifestations occur as progressive, irreversible obstruction (bronchiolitis obliterans) and less frequently lymphocytic alveolitis resulting in interstitial fibrosis or bronchiolitis obliterans organizing pneumonia (BOOP) (see Chap. 52).

Since the onset of pulmonary symptoms may not be symptomatic and obstruction may be irreversible, regular evaluations of a serial pulmonary function test (PFT) with body plethysmography (from the age of 4–6 years on) and diffusion capacity (usually from 8–10 years of age on) are required in asymptomatic patients.

While interstitial fibrosis is well known after lung transplant (restrictive allograft syndrome), prospective data after allogeneic HSCT are lacking, but case reports indicate that restrictive immune-mediated lung disease after allo-HSCT may occur.

Patients require follow-up by a pediatric experienced pulmonologist. Of note, the possible overlap of (1) myopathy/hypotrophy of the respiratory muscles (glucocorticoid induced, ± central obesity, and/or physical inactivity), (2) restriction of the chest wall in the context of dermal sclerosis, and (3) unproportional chest growth after TBI and/or local irradiation may contribute with a restrictive ventilator dysfunction leading to a mixed picture.

Finally, a thorough diagnostic evaluation includes a lung CT scan and a BAL to rule out viral, bacterial, fungal, and mycobacterial infections.

Coexisting IgA deficiency and chronic sinusitis or sinubronchial syndrome should be considered in the diagnostic workup (Hildebrandt et al. 2011).

44.2.8 Joints and Fasciae

cGVHD-associated fasciitis (diagnostic for cGVHD) can result in restricted mobility of joints. This can also be caused by deep cutaneous sclerosis. Moreover, rheumatoid complaints may be associated with cGVHD. In children myositis, muscle weakness, cramping, edema, and pain are quite common. However, iatrogenic glucocorticoid-induced myopathy may overlap with fasciitis. Range-of-motion (ROM) examinations are recommended at baseline and at serial intervals with the P-ROM scale providing an easy-to-apply tool. (There is a pediatric adaption, ped P-ROM; see addendum).

44.3 Diagnosis

cGVHD is diagnosed on the basis of cGVHD symptoms of eight organs, laboratory values (for hepatic manifestations), and PFTs. Each organ is graded between 0 and 3. The overall severity of cGVHD is classified as mild, moderate, or severe based on this organ-specific grading (number of organs and severity). Overall severity is calculated on the basis of the number of organs affected and the severity of their involvement. Only in case that functional involvement is solely due to none GVHD causes the impairment is not scored (Jagasia et al. 2015). Biomarkers of cGVHD are currently explored but require validation before clinical use.

44.3.1 Organ Grading of cGVHD for Adults and Children (See Annex 1 and Addendum)

44.3.2 Grading of Overall Severity of cGVHD (Jagasia et al. 2015)

Overall severity	Mild	Moderate	Severe
Number of involved organs	1–2	≥3	<u>>3</u>
Severity of involved organs	Mild (excluding lung)	Mild- moderate (lung only mild)	Severe (lung moderate or severe)

If diagnostic symptoms of cGVHD are absent, histological confirmation of diagnosis may be required. This may be particularly the case in gastrointestinal, nonspecific cutaneous, hepatic, and pulmonary manifestations to rule out toxic or infectious causes or comorbidity. Clinicopathologic series indicate a significant risk for inappropriate diagnosis and subsequent treatment if diagnosis has been made solely by clinical manifestations (and lacking diagnostic symptoms) without histological confirmation.

44.4 Treatment

44.4.1 First-Line Therapy

First-line treatment (see Table 44.1) consists of steroids given alone or in combination with CNI and is based on randomized trials.

As *mild cGVHD* does not impair organ function, the use of topical IS (topical steroids, topical CNI, or phototherapy) should be considered. If this is impossible, PRD treatment at an initial dose of 0.5–1 mg/kg body weight/day is recommended. Topical IS can be used in addition to systemic IS, to improve efficacy, or to reduce systemic IS, but lack systemic efficacy.

For *moderate or severe cGVHD*, systemic treatment with PRD or methylPRD at an initial dose of 1 mg/kg body weight/day should be used. In individual cases lower doses of 0.5–1 mg/kg may be used (Jacobsohn 2010). The combination of steroids with a CNI (CSA or TAC) is particularly worth considering for severe cGVHD. Rituximab has been explored in first-line treatment of cGVHD in combination with steroids and CNI demonstrating an increased response rate on the expense of an increased risk for late infectious complications and delayed B-cell recovery. Currently, ECP and ibrutinib are evaluated in first-line treatment of cGVHD within randomized clinical trials.

As cGVHD often takes time to respond to IS treatment, response should not be assessed until at least 8 weeks have elapsed or until 3–6 months have elapsed in the presence of deep cutaneous sclerosis. Long-term IS treatment lasting at least 3–6 months is often required. Dose reduction of IS agents should be performed stepwise.

	Recom	mendation	Side effects in >25%	Response	
Drug	Grade	Evidence	patients	rate	Comment
Steroids	A	I	Osteoporosis, osteonecrosis, diabetes mellitus	~30–50% CR	Main drug; strategies to reduce use due to SEs very important
CNI + steroids	C-1	II	Renal toxicity, hypertension	~30–50% RC	Reduces steroid use, reduced incidence of osteonecrosis
Rituximab + steroids/CNI	C-1	III-1 ¹²	Increased risk for late infectious complications	~75%	Randomized data are lacking
MMF + CNI/ steroids	D	II	GI complaints, infections		No increased efficacy compared to CNI and steroids, increased risk of relapse of malignancy
Azathioprine	D	II	Cytopenia, risk of infection		Increased mortality
Thalidomide	D	II	Neurotoxicity, drowsiness, constination		Very little effect in first-line therapy

Table 44.1 First-line treatment of cGVHD

Adapted from Wolff et al. (2011), A: should always be used; C-1: use in first-line therapy justified, D: moderate evidence of lack of efficacy or unacceptably high risks, should generally not be offered, I: evidence from ≥ 1 properly randomized, controlled trials, II: evidence from more than one well-planned non-randomized clinical trial, from cohort or case-controlled, analytic studies (preferably at several sites), III-1: only one non-controlled study, III-2: only one retrospective, non-controlled study or retrospective evaluation. (Evidence and recommendations graded according to the 2005 NIH Consensus), SE side effect, NIH US National Institutes of Health, MMF mycophenolate mofetil

Depending on the patient population, first-line therapy achieves complete remission of cGVHD in approximately 20% (adults) to 50% (children) of cases. If symptoms progress during the first 4 weeks of first-line therapy or there is no improvement in symptoms within 8–12 weeks, second-line therapy should be initiated.

44.4.2 Topical Therapy and Supportive Care

In principle, there is no difference between cGVHD treatment for children and adults. However, longterm steroid therapy in children causes major side effects in terms of growth, bone density, osteonecrosis, and organ development, making agents that reduce steroid use, entailing the use of topical drugs, particularly important. Age-based ancillary supportive care is essential in the management of pediatric cGVHD with the chance of sparing systemic therapy, often supported by highly compliant parents and/or family members as caregivers (Carpenter et al. 2015). In small children, the risk of systemic effects of topical steroid and CNI treatment must be considered. cGVHD is by itself remarkably immunosuppressive intensified by its treatment (especially high-dose corticosteroids)

leading to a high risk for infections: (a) for viral reactivation like CMV, ADV, and EBV and (b) for fungal infection like candida and aspergillosis. Functional asplenia with occurrence of Howell-Jolly bodies and a higher incidence of pneumococcal sepsis has to be considered also. Breakdown of skin and mucosal barriers adds to this risk.

Revaccinations (see Chap. 29) with inactivated vaccines are strongly recommended after consolidation of cGVHD (Hilgendorf et al. 2011). Live vaccines should be avoided in this patient population. Ursodeoxycholic acid reduced liver GVHD and improved survival (Ruutu et al. 2014). Supplemental IVIG replacement is recommended in cGVHD patients with IgG <400 mg/dL or recurrent infections which is of special importance in children but does also apply to adults. In case of long-term substitution or the history of anaphylactic reactions, we prefer to substitute subcutaneously.

44.4.3 Second-Line Therapy

While first-line therapy is based on randomized trials, second-line therapy mostly is based on phase II trials, and retrospective analyses are available (see Table 44.2). In addition, because

 Table 44.2
 Second-line treatment of cGVHD

	Recom	mendation		Side effects in >25%	
Drug	Grade	Evidence	Response rate	of patients	Comments
Steroids	В	III-1	n.a.	Osteoporosis, osteonecrosis, diabetes mellitus	Main drug, strategies to reduce use due to SEs very important
Ibrutinib	C-1	III-1	~50–75% ~16–25% CR	Bruising, diarrhea, infections	FDA approved in second-line treatment of cGVHD
Photophereses	C-1	II	~60–70% ~30% CR	Infections of the CVC (if applicable)	Venous access required, steroid-saving effect, good tolerability
mTOR-inh (sirolimus, everolimus)	C-1	III-1	~60% ~20% CR	TMA, hyperlipidemia, cytopenia	Increased risk of TMA when combined with CNI, regular blood levels required
MMF	C-1	III-1	~50% ~10% CR	GI SEs, risk of infection (viral) and increased risk of relapse	Steroid sparing activity
CNI	C-1	III-1	n.a.	Renal toxicity, hypertension	Reduces steroid use, regular blood levels required
MTX	C-2	III-1	~50% ~10–20% CR	Cytopenia	Best results in mucocutaneous cGVHD, reduces steroid use, contraindicated in the presence of pleural effusions or ascites
IL-2	C-2	III-1	~65% (only PR)	Fever, malaise, and fatigue	Applied in sclerodermoid skin disease
Ruxolitinib	C-2	III-1	n.a. (retrospective analysis)	Increased risk for viral reactivation, bacterial infection, hepatotoxicity	Prospective data pending
Bortezomib	C-2	III-1	n.a. for second-line Tx	Cytopenia, neuropathy	Trial was performed in first-line treatment
High-dose steroids	C-2	III-2	50–75% (only PR)	Infections	Rapid control of cGVHD
Total nodal irradiation	C-2	III-2	~50% ~25% CR	Cytopenia	Best results for fasciitis and mucocutaneous cGVHD
Hydroxychloroquine	C-2	III-2	~25% ~10% CR	GI side effects	Best results for mucocutaneous and hepatic cGVHD
Pentostatin	C-2	II	~50% ~10% CR	Cytopenia, risk of infection	Best results in children
Rituximab	C-2	П	~50% ~10% CR	Risk of infection	Effective in manifestations associated with autoAb and sclerodermoid cutaneous involvement
Imatinib	C-2	III-1	~50% ~20% CR	Fluid retention	Efficacy demonstrated mainly in sclerodermoid cGVHD and bronchiolitis obliterans
Thalidomide	C-3	II	~20–30% (only PR)	Neurotoxicity, drowsiness, constipation	Treatment for simultaneous cGVHD and recurrent multiple myeloma

(continued)

Table 44.2 (continued)

	Recommendation			Side effects in >25%	
Drug	Grade	Evidence	Response rate	of patients	Comments
Azathioprine	C-3	III-1	n.a.	Cytopenia, risk of infection, secondary malignancies	Increased risk of malignant disease of the oral mucosa
Retinoids	C-3	III-2	~60% (only PR)	Skin toxicity, hyperlipidemia	Effective in sclerodermoid cutaneous involvement
Abatacept	C-3	III-2	~40%		Effective in mucocutaneous and pulmonary involvement
Regulatory T cells	C-4				Currently explored in several clinical trials
Mesenchymal stem cells	C-4	III-2	n.a.		Repetitive application required
Alemtuzumab	C-4	III-3	n.a.	Infectious risks	Last resort for refractory cGVHD
Etanercept	C-4	III-3	n.a.	Infectious risks	May be used to treat mixed acute and chronic GVHD or pulmonary or GI manifestations of cGVHD

Adapted from Wolff et al. (2011), B: should generally be used, C-1: use in second-line therapy justified, C-2: use after failure of second-line therapy justified, C-3: should only be used in specific circumstances, due to unfavorable risk profile, C-4: experimental, should only be used in clinical trials and individual cases, II: evidence from >1 well-designed clinical trial without randomization, from cohort or case-controlled analytic studies (preferable from >1 center) or from multiple time series, III-1: several reports from retrospective evaluations or small uncontrolled clinical trials, III-2: only one report from small uncontrolled clinical trial or retrospective evaluations, III-3: only case reports available, SE: side effect, n.a.: not available

the data on disease severity and patient populations are very heterogeneous (in terms of age, conditioning, and stem cell source), the published response rates cannot be fully extrapolated to the majority of patients currently treated for cGVHD. Moreover, many substances have been used almost exclusively in combination with steroids.

In general, no more than three IS agents should be combined, as combinations of more drugs often does not lead to improved efficacy but results in a significantly increased risk of side effects and infections. Because of the substantial toxicity of long-term steroid treatment, strategies for dose reduction are very important. Since no predictors of response for a single agent in individual patients are yet available, the choice of

agent depends mainly on side effect profiles and patients' medical history. The response rates for specific agents range between 20% and 70% (photopheresis).

Certain drugs such as imatinib and retinoids are recommended only for manifestations associated with sclerosis (bronchiolitis obliterans [imatinib], sclerodermoid cutaneous alterations [retinoids, imatinib]), because of their specific mechanisms of action.

Response is assessed as for first-line therapy. Administration of drugs that have been shown to be ineffective should be stopped. As a rule, drugs shown to be ineffective should be tapered off stepwise with no more than one drug to be changed at a time in order to be able to evaluate their efficacy.

Appendix 1

Annex 1 - Organ Scoring of Chronic GVHD

	SCORE A	SCORE 1	SCORE 2	SCORE 2		
DEDECOMANCE COORE.	SCORE 0	SCORE 1	SCORE 2	SCORE 3		
PERFORMANCE SCORE:	☐ Asymptomatic	☐ Symptomatic,	☐ Symptomatic,	☐ Symptomatic,		
	and fully active	fully ambulatory,	ambulatory,	limited self-		
KPS ECOG LPS	(ECOG 0; KPS or LPS 100%)	restricted only in	capable of self-care, >50% of waking	care, >50% of		
	LF3 100 /8)	physically strenuous	hours out of bed	waking hours in bed (ECOG 3-4,		
		activity (ECOG 1,	(ECOG 2, KPS or LPS	KPS or LPS		
		KPS or LPS 80-	60-70%)	<60%)		
		90%)	00-7076)	100 /6)		
SKIN†		3070)				
O.t.i.vi						
SCORE %BSA						
GVHD features to be	□ No BSA involved	☐ 1-18% BSA	☐ 19-50% BSA	□ >50% BSA		
scored						
by BSA:						
Check all that applies:						
☐ Maculopapular						
rash/erythema						
☐ Lichen planus-like						
features						
☐ Sclerotic features						
☐ Papulosquamous						
lesions or ichthyosis						
☐ Keratosis pilaris-like						
GVHD						
SKIN FEATURES				Check all that		
SCORE:	☐ No sclerotic		☐ Superficial	applies:		
	features		sclerotic features	☐ Deep sclerotic		
			"not hidebound"	features		
			(able to pinch)	☐ "Hidebound"		
				(unable to pinch)		
				☐ Impaired mobility		
				☐ Ulceration		
Other skin GVHD feature	es (NOT scored by BSA)	_				
Check all that applies:						
☐ Hyperpigmentation						
☐ Hypopigmentation						
☐ Poikiloderma						
☐ Severe or generalized	pruritus					
☐ Hair involvement						
☐ Nail involvement						
☐ Abnormality present b	ut explained entirely by no	n-GVHD documented cau	ise (specify):			
MOUTH	☐ No symptoms	☐ Mild symptoms	☐ Moderate	☐ Severe symptoms		
Lichen planus-like		with disease signs	symptoms with	with disease signs		
features present:		but not limiting oral	disease signs with	on examination		
☐ Yes		intake significantly	partial limitation of	with major		
□No			oral intake	limitation of oral		
				intake		
☐ Abnormality present but explained entirely by non-GVHD documented cause (specify):						

Annex 1 - Organ Scoring of Chronic GVHD (continued)

	SCORE 0	SCORE 1	SCORE 2	SCORE 3			
EYES Keratoconjunctivitis sicca (KCS) confirmed by Ophthalmologist: Yes No Not examined	□ No symptoms	☐ Mild dry eye symptoms not affecting ADL (requirement of lubricant eye drops ≤3 x per day)	Moderate dry eye symptoms partially affecting ADL (requiring lubricant eye drops > 3 x per day or punctal plugs), WITHOUT new vision impairment due to KCS	Severe dry eye symptoms significantly affecting ADL (special eyeware to relieve pain) OR unable to work because of ocular symptoms OR loss of vision due to KCS			
☐ Abnormality present b	ut explained entirely by no	on-GVHD documented cau	ıse (specify):				
GI TRACT Check all that applies: Esophageal web/ proximal stricture or ring Dysphagia Anorexia Nausea Vomiting Diarrhea Weight loss* Failure to thrive	□ No symptoms	☐ Symptoms without significant weight loss* (<5%)	☐ Symptoms associated with mild to moderate weight loss* (5-15%) OR moderate diarrhea without significant interference of daily living	Symptoms associated with significant weight loss* >15%, requires nutritional supplement for most calorie needs OR esophageal dilation OR severe diarrhea with significant interference of daily living			
☐ Abnormality present b	ut explained entirely by no	on-GVHD documented cau	use (specify):				
LIVER	☐ Normal total bilirubin and ALT or AP <3 x ULN ut explained entirely by no	☐ Normal total bilirubin with ALT ≥3 to 5 x ULN or AP > 3 x ULN on-GVHD documented cau	☐ Elevated total bilirubin but ≤3 mg/dL or ALT > 5 ULN use (specify):	☐ Elevated total bilirubin > 3 mg/dL			
LUNGS** Symptoms score:	□ No symptoms	☐ Mild symptoms (shortness of breath after climbing one flight of steps)	☐ Moderate symptoms (shortness of breath after walking on flat ground)	☐ Severe symptoms (shortness of breath at rest; requiring 0₂)			
Lung score: FEV1	☐ FEV1≥80%	☐ FEV1 60-79	☐ FEV1 40-59%	☐ FEV1 ≤39%			
Pulmonary function tests ☐ Not performed	Pulmonary function tests ☐ Not performed						
☐ Abnormality present but explained entirely by non-GVHD documented cause (specify):							

Annex 1. Organ scoring of chronic GVHD (continued)

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	SCORE	0	SCORE 1		SCORE 2	SCORE 3
P-ROM score (see below)	□ No symptom	is	☐ Mild tightness of arms or legs, normal or mild decreased range	or le	ntness of arms egs OR joint tractures, hema	☐ Contractures WITH significant decrease of ROM AND significant
Shoulder (1-7):			of motion (ROM)	,	ight due to	limitation of ADL
Elbow (1-7):			AND not		iitis,	(unable to tie
Wrist/finger (1-7):			affecting ADL		derate	shoes, button shirts, dress
Ankle (1-4):				dec	rease ROM o mild to	self etc.)
					derate	
					ation of ADL	
☐ Abnormality present	t but explained e	ntirely i	by non-GVHD documer	ited cau	se (specity): 	
GENITAL TRACT	☐ No signs		☐ Mild signs‡ and	□ Mod	derate signs‡	☐ Severe signs‡ with
(See Supplemental			females with or	and	may have	or without
<u>figure</u> ‡)			without	sym	ptoms* with	symptoms
Check all that applies			discomfort on	disc	comfort on	
□ Not examined			exam	exa	m	
Currently sexually						
active						
☐ Yes						
□ No						
☐ Abnormality present	t but explained e	ntirely I	by non-GVHD documer	ited cau	se (specify):	
	severity (0-3) b		lications related to ch in its functional impac			
☐ Ascites (serositis)	_	☐ Mya	sthenia Gravis			
☐ Pericardial Effusion_		□ Peri	pheral Neuropathy		☐ Eosinophilia	> 500µl
☐ Pleural Effusion(s)_		□ Poly	myositis		☐ Platelets <1	00,000/μl
☐ Nephrotic syndrome)	□ Wei	□ Weight loss* without GI		☐ Others (specify):	
		sympt	mptoms			
Overall GVHD Severity (Opinion of the evaluator)	□ No GVHD		□ Mild	□ м	oderate	□ Severe
Photographic Range of Motion (P-ROM)						
	Shoulder	1 (Worst)	2 3 4 5 T T T	6 7(No	mat	
	Elbow	1 (Morat)	2 3 4 5	6 700	ma)	
	Wrist/finger	1 (Marri	大大大	6 7 (No		
	Ankle	1	1 1 1			

Adapted from Jagasia, 2015.

- † Skin scoring should use both percentage of BSA involved by disease signs <u>and</u> the cutaneous features scales. When a discrepancy exists between the percentage of total body surface (BSA) score and the skin feature score, OR if superficial sclerotic features are present (Score 2), but there is impaired mobility or ulceration (Score 3), the higher level should be used for the final skin scoring.
- * Weight loss within 3 months.
- ** Lung scoring should be performed using both the symptoms and FEV1 scores whenever possible. FEV1 should be used in the final lung scoring where there is discrepancy between symptoms and FEV1 scores.

 Abbreviations: ECOG (Eastern Cooperative Oncology Group), KPS (Karnofsky Performance Status), LPS (Lansky Performance Status); BSA (body surface area); ADL (activities of daily living); LFTs (liver function tests); AP (alkaline phosphatase); ALT (alanine aminotransferase); NUL (normal upper limit).
- ‡ To be completed by specialist or trained medical providers (see Supplemental Figure).

Appendix 2

Diagnosis and staging cGVHD in children

Jagasia et al BBMT 2015

pediatric adaptation A. Lawitschka 11/2015

patient name

date:

□ both

patient name

▶ please score/check the worst manifestation

► diagnostic features are marked **bold**

classification:actual

onset type ONLY at diagn.:

□ feat. of acute GVHD□ feat.of classic cGVHD

□ quiescent□ progressive

□ de novo

symptoms/features	Score 0	Score1	Score 2	Score 3
KPS/LPS: %	□ asymptomatic and fully active (KPS/LPS 100%)	sympt., fully amb., restricted only in physically strenous activity (KPS/LPS 80-90%)	sympt., amb., capable of self-care, >50% of waking hours out of bed (KPS/LPS 60-70%)	usympt., limited self-care >50% of waking hours in bed (KPS/LPS < 60%)
SKIN				
eat. scored by BSA:	no BSA involved	1-18% BSA	19-50% BSA	> 50% BSA
maculopapular rash/erythema				
lichen planus-like features				
sclerotic features:				0
□ lichen sclerosus-like				
□ morphea-like				
papulosquamous lesions				
ichthyosis				
keratosis pilaris-like GVHD				0
Feat. not scored by BSA: hyperpigmentation hypopigmentation/depigmentation poikiloderma severe pruritus hair involvement nail involvement sweat impairment abnormality present but explained entirely by non-GVHD cause (specify): feature decisive for diagnosis /scoring:				%BSA: child: head front/back 9 / 9 back 18, chest 18, arm left 9, arm right 9 leg left 13,5, leg right 11 adult: head front/back 4,5 / 4,5 back 18, chest 18 arm left 9, arm right 9 leg left 18, leg right 18 palm: 1,5
sclerotic features:	□ no sclerotic	w v	□ superficial	□ deep sclerotic features
	features		sclerotic features	"hidebound" (unable to pinch)

"not hidebound"

□ impaired mobility

(able to pinch) ulceration

MOUTH	M	Οl	JΤ	Ή
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□ erythema □ mild sympt with moderate sympt. $\ \square$ no symptoms severe sympt. with □ lichen planus-like features with disease signs disease signs but disease signs on examination not limiting oral with partial limitation with major limitation □ hyperkeratot. plaques intake significantly of oral intake □ mucoceles □ pseudomembranes of oral intake □ ulcers □ mucosal atrophy

□ abnormality present but explained entirely by non-GVHD cause (specify):

► feature decisive for diagnosis /scoring:

Appendix 2 - Diagnosis and staging cGVHD in children (continued)

symptoms/featu	ires	Score 0	Score1	Score 2	Score 3
EYES					
□ photophobia		□ no symptoms	☐ mild dry eye sympt. not affecting ADL (requirement of lubricant eye drops ≤ 3 x per day)	moderate dry eye sympt. partially affecting ADL (lubricant eye drops >3 x/d or punctual plugs) without new vision impairement due to KCS	□ severe dry eye sympt. significantly affecting ADL (special eyeware to relieve pain) unable to work because of ocular sympt or loss of vision due to KC
abnormality present	but explained entire	y by non-GVHD cause (snecify):	imparement due to NOO	
	r diagnosis /scoring:	y by non-dvrib cause (specify).		
GI TRACT	3				
esophageal web/		□ no symptoms	□ symptoms without	□ sympt. associated with	□ symptoms associated with
prox stricture or i	ring		significant weight	mild to moderate	significant weight loss (> 15%)
□ dysphagia	□ abdominal pain		loss (5%)	weight loss (5-15%)	requires nutritional supplement for
anorexia a	□ failure to thrive			or moderate diarrhea	most calorie needs or
nausea	□ vomiting			without significant	esophageal dilatation or
□ diarrhea	□ weight loss ≥ 5%			interference with	severe diarrhea with
				daily living	signif. Interference with daily livir
abnormality present	but explained entire	y by non-GVHD cause (specify):		height:
 feature decisive for 	diagnosis /scoring:				weight:
LIVER					
hepatic pattern		normal total bili	□ normal total bili	 elevated total bili 	□ elevated total bili > 3 mg/dl
Bili: AST:_	ALT:	and ALT or AP	with ALT ≥ 3-5x ULN	but ≤ 3 mg/dl or	
GGT: AP: _		< 3 ULN	or AP ≥ 3 x ULN	ALT > 5 ULN	
	•	y by non-GVHD cause (specify):		
 feature decisive for 	diagnosis /scoring:				
LUNGS					
		no symptoms	□ mild symptoms	□ moderate symptoms	□ severe symptoms
	MEF50: %	FEV1 ≥ 80%	(shortness of breath	(shortness of breath	(shortness of breath at rest;
	MEF75: %		after climbing one	after walking on	requiring O2)
	□ RV/TLC > 120%		flight of steps)	flat ground)	FEV1 ≤ 39%
CT:			FEV1 60-79%	FEV1 40-59%	
	•	y by non-GVHD cause (specify):		
feature decisive for	-				
JOINTS AND					
ped P-ROM score (s	*	□ no symptoms	□ mild tightness,	□ tightness or joint	□ contractures, fasciitis
	□ fasciitis		normal or mild ↓ of	contractures, fasciitis,	significant ↓ of ROM,
□ muscle cramps	□ athralgia		range of motion (ROM)	moderate ↓ of ROM,	significant ↓ of ADL
		0.415	not affecting ADL	mild - moderate ↓ of ADL	
		y by non-GVHD cause (specify):		
 feature decisive for GENITAL TF 					
erosions, fissures		□ no signs	□ mild signs	□ moderate signs	□ severe signs with or without
		1 IIO SIGIIS	□ IIIIu sigris	□ moderate signs	-
□ lichen planus-like □ lichen sclerosus-li					symptoms
ilicnen scierosus-ii ⊒ labial/ vaginal scai					
		y by non-GVHD cause (enecifu):		
 abnormality present feature decisive for 	•	y by non-GvnD cause (apeuty).		
. Julius decisive IOI	alagricolo /occirily.				
	it.				
Overall GVHD se	verity				
Overall GVHD se no cGVHD	verity				

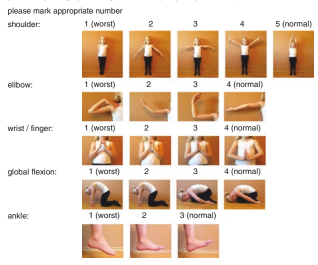
□ moderate: ≥3 organ with max score 1 or max. score of 2 in any affected organ, lung score max 1

 $\hfill \square$ severe: score 3 in any affected organ, lung score 2-3

Appendix 2 - Diagnosis and staging cGVHD in children (continued)

Other indicators, clinical featur	biopsy:		
check all that apply and assign a	organ:		
□ ascites (serositis)	☐ myasthenia gravis	☐ eosinophilia >500 /ul	GVHD confirmed?
☐ pericardial effusion	☐ peripheral neuropathy	□ platelets <100 000/ul	
□ pleural effusion	□ polymyositis	☐ hypo/hyperglobulinemia	
□ nephrotic syndrome	☐ weight loss >5% without GI sympt	□ auto-antibodies	
□ others (specify)	☐ diabetes		

pediatric photographic range of motion (adapted ped P-ROM):



Appendix 3

Genital Tract GVHD As	sessment and Scoring	Form		
Name:		Date of birth:		
Assessment date:				
	SCORE 0	SCORE 1	SCORE 2	SCORE 3
GENITAL TRACT (male or female)	☐ No signs	Mild signs and females may have symptoms* WITH discomfort on exam	Moderate signs and may have symptoms* with discomfort on exam	Severe signs with or without symptoms*
Currently sexually activ	e:	I.	I.	
Yes No				
Check all signs that a	pplies:			
Abnormality preser Abnormality though * Genital symptoms are infection.	te features emale) utination (female) emale) arring/ stenosis (male) arring/ stenosis (male) arring/ to repent to repent to repent to represent GVHD PL	resent GVHD (specify cau <u>USoth</u> er causes(specify cau d can represent premature	ause): e gonadal failure or geni	
as follows:		nation may be performed		
(Skene's and Bartho gentle touch of a qti	plin's), labia minora and r p is classified as discom	for the above signs. Toucl najora gently with a qtip. V fort on examination. Palpa owing or other signs of vag	ulvar pain elicited by the te the vaginal walls with	e
,		hether qtip palpation or ge oman experiences during i		
Female genitalia: Seven 1) Mild (any of the follow	, ,	mucosal surfaces, vulvar l	lichen-planus or vulvar li	chen-sclerosis.
2) Moderate (any of the f	ollowing); erosive inflam	matory changes of the vulv	var mucosa, fissures in v	vulvar folds.
, , ,	s vaginal banding, vagina	oral hood agglutination, fib al shortening, synechia, de	· ·	
-		n planus-like or lichen scle	rosis-like features and	
phymosis or urethral sca 1) Mild: lichen planus-like	=	y of signs:		
Moderate: lichen scler		erate erythema:		
3) Severe: phimosis or u		rate orythoma,		
Biopsy obtained: Yes	No Site biopsied:	GVHD conf	irmed by histology:	Yes No
Change from previous evalu	uation: No prior or	current GVHD Improv	red Stable	Worse N/A (baseline)
Completed by (spell out	t name):			

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