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Glucocorticoid receptor action in beneficial and side effects of steroid therapy: lessons from conditional knockout mice

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Summary

Glucocorticoids (GCs) are potent immune suppressive drugs with unfortunately severe side effects. Different molecular modes of actions of the GC receptor (GR) have been identified. Transcriptional transactivation by binding of a dimerized GR protein complex to the promoter of GC regulated genes or interference with activity of pro-inflammatory transcription factors by GR monomers are considered as the two major mechanisms. It has been hypothesized that selective GR agonists (SEGRAs) addressing dimer-independent function would reveal potent steroid therapeutic activity with reduced side effects. Recent studies of a mouse knock-in strain with a dimerization-deficient GR demonstrate that some inflammatory processes can be suppressed by GCs, while others can not. Also side effects of GCs occur in these mice. Thus, depending on the process that is treated, SEGRA could be therapeutically more or less effective and not all side effects of steroid therapy may be reduced.
Introduction

When cortisol related substances are prescribed to treat an allergic skin reaction, or a chronic ulcerative disease, medical doctors as well as patients are left with a feeling that the “evil is combated by the devil”. Why are these powerful anti-inflammatory drugs so double sided as the schizophrenic entities living in one person called Dr. Jekyll and Mr. Hyde? Are there possibilities to strengthen Dr. Jekyll and to weaken Mr. Hyde’s properties of cortisone like substances? Obviously a better understanding of the molecular nature of these factors and the contribution to the phenotypes of both gentlemen will allow us to speculate, if intelligent drugs will prevent Mr. Hyde in steroid therapy.

Glucocorticoids (GCs), such the synthetic drugs prednisolone, betamethasone, or the potent compound dexamethasone, belong to the class of steroids, which bind to the glucocorticoid receptor (GR). The endogenous ligands of the GR, cortisol, corticosterone and aldosterone – the latter at least acting glucocorticoid like in the brain – are synthesized in the adrenal cortex. Their production is diurnal in a pulsatile manner. The release of GCs is regulated by a hierarchy of endocrine organs The hypothalamus secretes corticotropin-releasing hormone (CRH) controlling the pituitary, which in turn triggers the release of adrenocorticotropic hormone (ACTH) that stimulates the steroid synthesis in the adrenal cortex. The hormonal network between these endocrine tissues is considered as the HPA (hypothalamus-pituitary-adrenal)-axis. The HPA axis is subject to intense negative feedback regulation by GCs levels themselves, finally leading to a circadian pulsatile behaviour of GC production. The input to the hypothalamus is either derived from the central circadian clock or – from psychological and physiological stress. The latter can be starvation, but also systemic inflammatory reactions.
As their name tells us, one of the primary functions of GCs is to increase serum glucose levels to allow rapid energy supply for the brain, which consumes preferably glucose. Anabolic processes in the liver achieve this by inducing gluconeogenesis and catabolic actions in peripheral organs by triggering protein degradation and increased lipolysis. One can envisage that these catabolic actions contribute much to the side effects, when GCs are applied at pharmacological doses. In particular the adverse effects manifest in numerous features such as brittle skin (Schoepe et al., 2006), muscle weaknesses, osteoporosis (Canalis and Delany, 2002), fat redistribution (Peeke and Chrousos, 1995), diabetes (Peeke and Chrousos, 1995), but also leads to neurodegeneration or suppressed neurogenesis in the brain (De Kloet et al., 1998).

Why are GCs still the therapeutic standard for the treatment of allergic diseases such as asthma, and of chronic inflammation such as rheumatoid arthritis and inflammatory bowel diseases? Dependent on the severity of the inflammatory disease the beneficial effects of GCs still balance out some of the adverse effects. In some cases non steroidal anti-inflammatory drugs (NSAIDs) can replace GC, whereas in other conditions GC therapy is combined with acute treatment of the side effects, e.g. to avoid GC induced osteoporosis by application of bisphosphonates (Canalis and Delany, 2002). However, there is still the ambitious goal in the pharmaceutical industry to produce steroidal analogs that avoid the side effects and maintain the therapeutic efficacy of steroid treatment.

The understanding of the molecular basis that underlie the therapeutic and side effects have been tremendously increased in the last years by the combination of molecular biology and the analysis of transgenic mice with cell type specific or function selective glucocorticoid receptor mutations. Based on these findings the first attempts of the design of selective GR agonists (SEGRA) have been performed and tested for
their therapeutic efficacy and side effects (Belvisi et al., 2001; Lin et al., 2002; Schacke et al., 2004; Vayssiere et al., 1997).

In this review we will summarize how molecular actions of the GC receptor (GR) contribute to certain therapeutic and certain adverse effects and finally we discuss how much hope lies in the pharmaceutical attempts to create potent anti-inflammatory GR ligands without side effects.

**Functional domains of the GR protein**

The gene NR3C1 encodes the nuclear receptor “glucocorticoid receptor”, which was cloned in the beginning of 1980ies in mammalian species (Evans, 1988). The GR was among the first described bona fide enhancer activating transcription factors in mammals (Miesfeld et al., 1984). The arrangement of functional domains of the GR protein – ligand-binding (LBD), DNA-binding (DBD) and transactivation domains - is common with most of the other nuclear receptors. The transactivation domain AF-1 is located in the N-terminus and its activity is subject to modulation by posttranscriptional modifications (Schaaf and Cidlowski, 2002), whereas the AF-2 transactivation domain is close to the C-terminus of the LBD in the C-terminus and changes to an active conformation after binding of a ligand. The DBD contains two so called Zn finger motifs. Each of this motif consists of two protein loops that are co-ordinated by a Zn atom. Both Zn fingers are followed by an amphipathic alpha helix (Luisi et al., 1991). The LBD is build up by a three-layer helical sandwich that embeds a hydrophobic pocket for ligand binding, which is followed by the C-terminal AF-2 containing helix 12 (Bledsoe et al., 2002; Shiau et al., 1998). In the absence of hormone the GR is localized in the cytoplasm, where it is in a complex with chaperone molecules, hsp90, hsp70, hsp40, the co-chaperone p23 and the immunophilins FKBP52 and Cyp40 (Pratt et al., 1996). Binding of the hormone leads
to disruption of the cytoplasmic complex and NLS sequences become unmasked and the GR molecules shuttles into the nucleus with the help of importin proteins (Freedman and Yamamoto, 2004).

The GR controls gene expression by several mechanisms. Some of them imply interaction with signal transduction components, such as PI3K, JNK, 14-3-3 proteins (Caelles et al., 1997; Kino et al., 2003; Limbourg and Liao, 2003) and take place partially in the cytoplasm. Some events even occur at the membrane as reviewed elsewhere in this issue [Buttgereit]. However, the best characterized actions of the GR are on the transcriptional level and employ two major mechanisms: i) binding to DNA and ii) interaction with other transcription factors by tethering mechanisms (Figure 1).

**Control of Gene expression by DNA binding of the GR**

The binding to DNA occurs on imperfect palindromic sequences present in the promoters and enhancer regions of GC responsive genes, designated as GC responsive elements (GRE). FRAP technology and chromatin-immunoprecipitation experiments exhibited a kinetic view for the DNA binding of the GR and other nuclear receptors. These experiments showed that the interaction with DNA is a dynamic process during which the GR and coactivator complexes cycle between DNA and nucleoplasm within seconds (McNally et al., 2000) or minutes (Metivier et al., 2003). The GR binds to GREs as a homodimer. Both half sites of the GRE are bound by one molecule of the dimer. The dimerization occurs via distinct hydrophobic motifs present in the ligand binding domain (Bledsoe et al., 2004) and is further stabilized by interaction in the DNA binding domain itself. Directly at the GRE half site the N-terminal of two Zn fingers contact the DNA whereas the second Zn finger contacts the partner molecule via an amino acid stretch called the dimerization box (Luisi et al., 1991). Thus, the dimerization is essential for binding of
the GR to GRE controlled genes. The mechanism of the binding of the GR to negative acting GREs (nGRE) so far identified only in a limited amount of genes, is less well understood, but seems also to require dimerization (Dostert and Heinzel, 2004).

Bound to the GRE the transactivation domains of the GR serve as docking platform for the recruitment of transcriptional coactivators that either remodel chromatin directly or facilitate the initiation of transcription. The AF-1 domain recruits in a ligand independent manner the multimeric chromatin-remodeling ATPase BRG1, a homologue of yeast SWI/SNF (Cairns et al., 1996; Yoshinaga et al., 1992) that in turn assemble the histone acetylases P/CAF and CBP/p300. The AF-2 domain upon binding of hormone recruits members of the p160 family of coactivators, which share a nuclear receptor interaction motif called the NR box (LxxLL). A number of these p160-factors, such as SRC-1, SRC-2/GRIP1/TIF2 and RAC3/ACTR/p/CIP possess histone acetylase (HAT) activity, leading to disruption of nucleosomes. Recently it could be demonstrated that an additional factor called STAMP seems to be part of the SRC-2/GRIP1/TIF2-GR complex and enhances its transcription-inducing activity (He and Simons, 2007). Furthermore additional chromatin-modifying enzymes such as the aforementioned CBP/p300 and the histone arginine methylase CARM1 (Chen et al., 1999; Ma et al., 2001) interact with the GR.

However, which co-activator/chromatin complexes are finally present on the promoters of genes depends on availability of coactivators in a certain cellular context, the abundance of other transcription factors competing for the same coactivators, packed stage of the chromatin and the phosphorylation status of the AF-1 domain. The mechanisms of positive regulation of gene transcription by the GR has been understood in detail only for a few well characterized promoters such as the MMTV, the tyrosine amino transferase gene (Schmid et al., 1987) and the
phosphoenolpyruvate carboxykinase (Hanson and Reshef, 1997). For a number of known GC induced genes, such as MAPK phosphatase-1 (Kassel et al., 2001) and GITR (Nocentini et al., 1997) knowledge of the precise mechanisms for up-regulation is poor. For the growing number of novel recognized GC induced genes by gene expression profiling (James et al., 2007; Rogatsky et al., 2003) a comprehensive mechanistic analysis is challenging. Recent advances with chromatin IP scanning of GC regulated promoters gave first insights of GR occupancy in sequences of some GC induced genes (Wang et al., 2004). The requirements of coactivator composition on individual promoters of GC regulated genes could be addressed by a combination of genome wide chromatin-immunoprecipitation approaches and traditional promoter sequence deletion analysis.

**Cross-talk of the GR with Transcription Factors**

Since 1990 a DNA binding independent mechanisms of GR action on transcription were discovered. Pro-inflammatory transcription factors such as AP-1 (Jonat et al., 1990; Schüle et al., 1990; Yang Yen et al., 1990), NF-κB (De Bosscher et al., 1997; Heck et al., 1997; Ray and Prefontaine, 1994; Scheinman et al., 1995b), IRF-3 (Ogawa et al., 2005; Reily et al., 2006) and STAT proteins (Stöcklin et al., 1996; Zhang et al., 1997) can be influenced in their activity to control transcription by GCs in the absence of a GRE, but in the presence of the binding sites of the interacting transcription factors. This protein-protein interaction is designated as “tethering mechanism” (Beato et al., 1995; De Bosscher et al., 2003). Mutational analysis of the GR revealed that the DNA binding domain was required for the interaction, even when the dimerization box of the second Zn-Finger was disrupted (Heck et al., 1994).
Whereas interaction of the GR with STAT5 leads to an increase of gene expression in cells (Stoecklin et al., 1997) and in mice (Tronche et al., 2004), for AP-1, NF-κB and IRF-3 a suppression of gene regulatory activity for most of the respective target genes was demonstrated (Göttlicher et al., 1998; Ogawa et al., 2005; Reichardt et al., 2001; Tuckermann et al., 1999). Numerous mechanisms have been proposed for the molecular nature of the inhibition of transcription factors by the GR. One involves a squelching model of limiting amounts of CBP/p300 (Kamei et al., 1996), which are competed out by the GR, and thereby attenuate AP-1 and NF-κB activity. However, De Boscher and colleagues demonstrated that repression of AP-1 and NF-κB is independent on CBP/p300 levels (De Bosscher et al., 2001). But the competing model seems to hold true for the inhibition of IRF-3 by the GR. In macrophages overexpression of the coactivator GRIP-1 abolished GC-inhibition of IRF-3 whereas knock down of GRIP-1 diminished IRF-3 activity to the same extend than GR activation by GC (Reily et al., 2006). NF-κB could be also inhibited indirectly by GC induced synthesis of its inhibitor IκBα (Auphan et al., 1995; Scheinman et al., 1995a) or by the GC induced leucine Zipper protein GILZ (D'Adamio et al., 1997), which was also demonstrated to inhibit AP-1 (Mittelstadt and Ashwell, 2001). As an additional mechanism GC induced expression of histone deacetylase 2 was shown to blunt NF-κB activity by recruitment of this HDAC to the p65-HAT-CBP complex (Ito et al., 2000). Direct interaction of the monomeric GR and pro-inflammatory transcription factors within a tethering complex that prevent their recruitment of co-activators (De Bosscher et al., 1997) or co-repressors or by interfering with interaction of RNA polymerase II (Nissen and Yamamoto, 2000) seems to be the best characterized mechanism of transrepression by the GR. The interaction between the GR and the transcription factors often requires integrator proteins such as the nuclear
isoform of the LIM-domain protein TRIP-6 (Kassel et al., 2004) and a recently identified factor STAMP (He and Simons, 2007).

**Disrupting dimerization-induced DNA-Binding in vivo: GR\textsuperscript{dim} mice**

In order to discriminate between the tethering mechanism and dimerization induced DNA binding of the GR the mutant GR-version described by Cato and colleagues (Heck et al., 1994) was knocked into mice (Reichardt et al., 1998). These GR\textsuperscript{dim} mice with a dimerization deficient receptor exhibit an absence of GRE regulated genes (Reichardt et al., 1998), but a preserved suppression of AP-1 (Tuckermann et al., 1999) and NF-κB-activity (Reichardt et al., 2001). Interestingly, these mice overcame the lethal lung phenotype of GR-knockout mice, stressing the importance of DNA binding independent activities of the GR for certain physiological processes (Reichardt et al., 1998). Furthermore irritant contact dermatitis by phorbol esters could be suppressed by GC in GR\textsuperscript{dim} mice (Reichardt et al., 2001). These findings lead to the promising concept that pharmaceuticals that would act as a SEGRA addressing the ability of the GR monomer by avoiding dimerization would promote Dr. Jekyll’s beneficial side and would omit Mr. Hyde’s deleterious side effects (Fig. 1). Is this promise realistic? We will come back to this question after we discussed in the remaining sections the tissue specific requirement of the GR and the involvement of dimerization induced DNA binding in therapeutic and side effects of GC.
Endogenous GCs in inflammatory responses: The example of Septic shock

In a variety of conditions the anti-inflammatory actions of endogenous GC are required to control massive immune responses. One example is sepsis. Sepsis is viewed as a complex dysregulation of inflammation arising when the host is unable to successfully defeat an infection. This exaggerated immune response that damages the organism, can lead to septic shock, frequently with lethal outcome. In rodents septic shock can be induced by a bolus injection of toll-like receptor (TLR) agonists such as lipopolysaccharides (LPS) and DNA (polyI:C), which bind to TLR4 and TLR3, respectively (Aderem and Ulevitch, 2000; Akira et al., 2006). In macrophages for example TLR4 activation leads to the recruitment of the intracellular adaptor molecules MyD88/IRAK/TRAF6 or TRIF/RIP-1 which causes the activation of MAP-kinases and IKKs (Akira et al., 2006). This results in the activation of the transcription factors AP-1, NF-κB, IRF-3 and others to trigger the induction of pro-inflammatory genes including cytokines associated with sepsis, such as TNF-α, IL-1 and IL-6. This finally converges on a systemic inflammatory response syndrome (SIRS). During this systemic inflammation GCs are synthesized in the adrenal gland in response to stress or systemic cytokine release following exposure to bacterial endotoxin (Yeager et al., 2004). Although treatment of septic shock with high GC doses in humans seems not to show a reduction of mortality (Meade, 2005), there are several lines of evidence that GCs indeed participate in control of this process. In patients that exhibit severe sepsis, adrenal insufficiency is accompanied. Furthermore, adrenalectomized rodents have difficulty in surviving mild septic shock reviewed in (Yeager et al., 2004). Recent studies with conditional GR mutant mice by us and others (Bhattacharyya et al., 2007), showed that the GR in myeloid cells is required to
survive sub-lethal LPS doses and cecal ligation and puncture (CLP). More important, to survive septic shock the regulation of GRE dependent gene expression is required, since knock-in mice with a dimerization deficient GR also do not survive lethal sepsis (Kleiman, unpublished). Thus, a transrepression mechanism, i.e. the down regulation of NF-κB, AP-1 or IRF-3 is not sufficient to lead to a successful management for sepsis. Genes that are induced by GC, such as the MAPK inhibitor DUSP-1 (MPK-1) have been hypothesized to be implicated in the immune suppressive effects of sepsis (Bhattacharyya et al., 2007). However, normal DUSP-1 protein levels in GR<sup>dim</sup> macrophages have been observed (Abraham et al., 2006), although GR<sup>dim</sup> animals do not survive sepsis. Which GC regulated genes could mediate the survival of a sub-lethal sepsis conditions? The answer of this question is complex, since numerous genes were identified to be regulated by GC in murine macrophages as revealed by gene expression profiling studies (Ogawa et al., 2005). A recent study of global gene expression in GC treated human monocytes (Ehrchen et al., 2007) revealed that GC not only suppress inflammatory functions of monocytes but rather shift them to an “anti-inflammatory” activated phenotype that includes enhanced phagocytic properties and migratory behaviour and a protection from apoptosis. These findings together with previous observations confirm that GC-treatment could stimulate monocytes/macrophages to optimize the clearance of pro-inflammatory complexes, dying neutrophils and finally to a resolution of the inflammation (Yona and Gordon, 2007).

Also the requirement of the GR in other cell types for the survival of sepsis cannot be excluded. The conditional GR mutant GR<sup>lymCre</sup> mouse lacks the GR not only in macrophages also in some other myeloid cell populations, in particular neutrophils. Whereas animals lacking the GR in T-cells show no difference in sepsis responses
animals that have no GC response in other compartments, such as endothelial cells have not been analyzed, yet. In summary, for the beneficial effect of GC in sepsis the presence of the GR at least in myeloid cells is required and dimerization induced DNA binding is needed.

_exogenous GC action in skin inflammation_

Inflammatory skin diseases, such as contact and atopic dermatitis are frequently prescribed with GCs with the long term risk of side effects, namely atrophy of the skin (Schoepe et al., 2006). The phorbol ester induced edema formation is a commonly used model for assessing unspecific and irritant skin inflammation (Gschwendt et al., 1984) and GCs and analogous compounds have been extensively tested for their anti-inflammatory action. For the suppression of this type of inflammation the dimerization of the GR is not required, since GR\textsuperscript{dim} mice show an efficient reduction of phorbol ester inflammatory response (Reichardt et al., 2001). In this model also function-selective GR ligands have been tested for their anti-suppressive efficacy (Schacke et al., 2004; Vayssiere et al., 1997). From the croton oil experiments one could conclude that dimerized-induced DNA binding can be omitted for suppression of inflammatory skin diseases. However, recent studies in contact hypersensitivity, a model for contact dermatitis and thus an inflammatory type of a different mechanism revealed that more than transrepression of the GR is required for suppression of inflammation (Tuckermann et al., 2007).

Contact dermatitis is represented by an experimental rodent model, contact hypersensitivity (CHS) and has been extensively studied over decades. Once seen as a rather simple allergic reaction independent on a humoral response, only dependent on cell-cell-interactions with three main players, antigen presenting cells, T cells and
effector cells, recently the picture has become more complex. The CHS reaction is divided into two phases, the sensitization phase and the challenge phase (Figure 2). The current view (Askenase, 2001) tells us that small molecular compounds called haptens when applied the first time to the body are covalently bound to epidermal proteins. This irritates keratinocytes, which subsequently activate antigen presenting cells (APCs). The APCs – notably dermal DCs triggered by cytokines such as TNF-α and IL1β – take up the hapten loaded proteins, migrate to draining lymph nodes and present the processed haptens to naïve T cells on MHC II molecules. In turn, T cells differentiate into antigen-specific Th1 and cytotoxic T cells and are designated as sensitized. Sensitized T cells proliferate, leave the local lymph node and patrol through the body. In the challenge phase, initiated by the re-exposure of the same hapten, activated endothelium allow the penetration of a first wave of lymphocytes, including the surveying sensitized T cells. These T cells become restimulated by APCs in the skin and subsequently release pro-inflammatory mediators, which trigger resident myeloid cells to secrete chemokines, finally leading to a fully inflammatory reaction including a second wave of leukocyte infiltration and edema. So there are numerous potential targets for GC action in this system to suppress the inflammatory response. We could show recently (Tuckermann et al., 2007), that anti-inflammatory action of GC is dispensable in keratinocytes, T-cells and presumably antigen presenting cells: Whereas conditional GR-knockout mice for Keratinocytes (GR^{K14Cre}) and T cells (GR^{LCKCre}) mount a perfect immune suppressive response by GCs, GR^{lysMCre}-mice, lacking the GR in macrophages and neutrophils were refractory to GC treatment. Thus, the GR in macrophages and neutrophils seems to be critical for steroid therapy. Interestingly, in contrast to irritant contact dermatitis mice with a dimerization defective GR (GR^{dim}) were largely resistant to GC treatment and
exhibited a persistant leukocyte infiltration after steroid therapy. Macrophages isolated from GRdim mice showed impaired suppression of IL-1β, MIP-2, MCP-1 and IP-10 by GC (Tuckermann et al., 2007), all cytokines that functionally counteracted GC effects on CHS. TNF-α showed only a minor effect. Thus, depending on the type of skin inflammation different cell types and different molecular mechanisms can be involved.

This has consequences for therapeutic strategies, for example one would expect that function selective GR ligands (SEGRAs) that address the transrepression activity with the hope to reduce atrophy of the skin (Schoepe et al., 2006) would be potent in the treatment of irritant dermatitis, but less in contact allergy.

**Side effects: GCs and Glucose homeostasis**

The most predominant side effects of GCs in the organism are of catabolic nature i.e. remodeling of tissue, such as the aforementioned skin, but also in bone leading to osteoporosis, redistribution of fat and insulin resistance leading to diabetes. These catabolic actions are due to the capability of glucocorticoids to enhance glucose levels by stimulating gluconeogenesis, which in part relies on the degradation of proteins and modulating fatty acid metabolism. The central role of GCs to maintain glucose levels is evident by the findings that patients deprived from GC have low glucose levels (Addison’s disease), patients with GC excess (Cushing syndrome) exhibit glucose intolerance (Andrews and Walker, 1999). The clinical features are corroborated by findings in in CRH knock-out mice devoid of GCs that fail to counteract hypoglycemia (Jacobson et al., 2006). The maintenance of glucose by GC could be explained by several potential mechanisms. First the induction of enzymes involved in gluconeogenesis in the liver could be crucial. Mice deficient for the GR in
hepatocytes (GR<sup>alcre</sup>) suffer from hypoglycemia after starvation and do not upregulate gluconeogenetic enzymes (Opherk et al., 2004). Prednisolone administration stimulates expression of the tyrosine aminotransferase gene in contrast to the SEGRA compound ZK216348 that fail to elevate both TAT mRNA and glucose levels in the blood (Schacke et al., 2004). Second, decreased glucose uptake in peripheral organs could be prevented by GC by inhibition of glucose transporter (GLUT4) translocation to the cell membrane (Horner et al., 1987), and stimulation of lipolysis by GC could counteract insulin mediated reduction of blood glucose levels (Andrews and Walker, 1999). Third, GC could prevent insulin production. Mice over expressing the GR in pancreatic beta-cells show impaired insulin production (Delaunay et al., 1997). If this directly occurs with normal GR levels in the excess of hormone has not been addressed so far. Fourth, GC-induced biosynthesis of ceramides, which inhibit Akt/PKB signaling could thereby lead to insulin resistance (Zierath, 2007). Recently, in DES1 knockout mice, which exhibit impaired ceramide synthesis, GC induced insulin resistance could be diminished (Holland et al., 2007). How much these potential mechanisms contribute to the GC-regulation of glucose levels and the GC-induced insulin resistance requires in depth analysis of GR cell type specific and GR function selective mutant mice.

**Side effects: GC actions on the skeleton**

GC long-term therapy has a strong impact on the skeletal system. Prolonged GC treatment of children with juvenile rheumatoid arthritis, chronic asthma and post renal transplantation strongly impairs longitudinal growth in children that receive steroid therapy (De Luca, 2006). Longitudinal growth in prepubertal vertebrates is occurring in the cartilaginous growth plate at the epiphysis of bones. In the growth plate chondrocytes undergo the serial events of proliferation, subsequent differentiation and
apoptosis which results in the replacement of cartilage by bone (Karsenty and Wagner, 2002). Apoptosis of terminal differentiated (hypertrophic) chondrocytes, osteoclastic resorption and the formation of bone matrix (mainly constituent collagen type I) by the osteoblasts lead to the replacement of cartilage by mineralized bone. GCs seem to influence the activity of the growth plate on several levels finally leading to reduced growth. First, GC impair proliferation of growth plate chondrocytes, by simultaneously enhancing apoptosis of hypertrophic chondrocytes (Smink et al., 2003b). Second, in vitro studies in cultured chondrocytes suggest that GCs induce apoptosis by activation of caspase-3 (Chrysis et al., 2003) and Bax (Mocetti et al., 2001), and decreased expression of Bcl-2 and Bcl-x (Mocetti et al., 2001). Third, GC seem to affect the GH/IGF-1 axis, which is important for proliferation and survival of chondrocytes on multiple levels. GC inhibit GH secretion in the pituitary (Devesa et al., 1995), decrease IGF-1 expression in the growth plate (Chrysis et al., 2003; Smink et al., 2003b), and down regulate GH receptor and GH binding protein (Gevers et al., 2002). In addition, GC impair IGF-1 signaling by inhibition of PI3K (Chrysis et al., 2005). Finally, the reduction of VEGF in chondrocytes points to a possible prevention of vascularogenesis leading to an impairment of growth by inhibition of mineralization (Koedam et al., 2002). The complex effects of GC on chondrocyte functions were recently reflected by results from expression profiling of GC treated chondrocytic micro mass cultures that recapitulate chondrocyte differentiation in vitro (James et al., 2005; James et al., 2007). GC treatment suppressed genes favoring proliferation such as growth factors, and enriched metabolic associated genes and genes encoding matrix genes, indicating that GC enhance chondrocyte differentiation. On the other hand factors promoting vascularization and mineralization were inhibited by GC (James et al., 2007). All
these changes of gene expression probably result in a reduced growth plate activity and a reduction of bone growth. However, studies with cartilage specific deletion of the GR are required to unequivocally dissect the GR function in cartilage cells versus effects on osteoblasts, osteoclasts and endothelial cells in the process of bone growth.

The other prominent side effect on the skeleton, GC induced osteoporosis seems now to become better understood. GC excess leads to a rapid bone loss that is followed by a longer lasting decline of bone formation and increasing the fracture risk to 50% for patients with more than 7.5 mg/kg prednisolone treatment (Van Staa et al., 2000). The effect of GC on bone homeostasis could be systemic or local. Systemic effects include a decreased calcium-absorption in the gut and decreased calcium-reabsorption in the kidney (Mazziotti et al., 2006). Lowering calcium-levels in the blood enhance PTH secretion from the parathyroid glands, which act on osteoblasts to stimulate osteoclast formation by inducing cortical bone loss. However, GC induced osteoporosis is accompanied by trabecular bone loss and reduced bone turnover. Hyperparathyroidism leads instead to increased bone turnover (Mazziotti et al., 2006), making it unlikely to be the cause. Furthermore the secretion of osteotropic hormones such as sex steroids and growth hormone secretion is attenuated (Mazziotti et al., 2006). Decrease of sex steroids by GC enhances bone loss and increases the risk for fractures. As for the actions in the growth plate inhibition of GH activity by decrease of IGF-I, GH receptor and IGF receptor 1 expression in osteoblasts as well as abrogating the release of GH from the pituitary could lead to a reduced bone strength (Delany et al., 2001; Devesa et al., 1995; Itagane et al., 1991; Smink et al., 2003a; Wehrenberg and Giustina, 1992).

In contrast to evidence for systemic actions, the direct effects on bone cells by GC leading to osteoporosis could be recently demonstrated with cell type specific
mutations of the GR. High dose GC treatment rapidly leads to impairment of bone formation that includes induction of apoptosis of osteoblasts and osteocytes in wild type mice (Weinstein et al., 1998). But apoptosis seems not to be the only mechanism. Osteoblast numbers are controlled by the canonical Wnt pathway (Hartmann, 2006) and recently GCs could be shown to impair Wnt signaling in osteosarcoma cell lines by direct binding of the GR to β-catenin (Takayama et al., 2006). Whereas low dose GC-treatment seems to stimulate osteoblast differentiation (Shalhoub et al., 1992), high dose GC treatment leads to inhibition of differentiation of primary mouse osteoblasts (Smith et al., 2000). The suppression of osteoblast function could be due to decreased expression of genes directly involved in bone formation, such as collagen 1 or runx2 (Pereira et al., 2001), but also due to antagonizing BMP pathways (Luppen et al., 2003). Also osteoclast-activity modulation by GCs participates in GC induced osteoporosis. GCs induce RANKL in osteoblasts, a potent stimulator of osteoclastogenesis (Hofbauer et al., 1999). GCs prolong longevity of osteoclasts in vivo (Jia et al., 2006; Kim et al., 2006). But bone degrading activity was also found to be suppressed by steroids (Kim et al., 2006). Furthermore in mice lacking the GR in osteoclasts a defect of bone formation activity by the osteoblasts was detected, linking osteoclast GC reactivity to osteoblast function (Kim et al., 2006). However, deleting the GR in osteoblasts themselves (Tuckermann, unpublished) also strongly ameliorated osteoblastic bone formation, while osteoclast numbers and activity remained intact. This is in agreement with studies of Manolagas and colleagues (Jia et al., 2006), which demonstrate that inhibition of GC action in osteoclasts in TRAP-11b-HSD2 transgenic mice still exhibit a decrease of osteoblast numbers and a decrease of bone formation. Thus, it seems so that inhibition of bone formation requires the cell autonomous GR of the osteoblasts, whereas actions of GCs in
osteclasts contribute also to the decrease of bone mineral density occurring in glucocorticoid induced osteoporosis. Finally we subjected GR\textsuperscript{dim} mice lacking the dimerization function of the receptor to high dose GC treatment and observed a similar inhibition of bone formation, osteoblast numbers and bone mineral density as in wild type mice (Tuckermann, unpublished). Interestingly, dimerization induced DNA binding is dispensible for this type of side effect and we have to hypothesize that protein-protein interactions with other transcription factors could be sufficient to mediate GC induced bone loss.

**Conclusion**

The analysis of the GR\textsuperscript{dim} mice in steroid therapy (Table 1) so far revealed that GC could successfully treat irritant dermatitis. GCs could efficiently suppress inflammatory regulators, such as TNF or MMP-1 and MMP-13 in these mutant mice, indicating that for therapeutic action of GC dimer-independent mechanisms of the GR are sufficient, such as tethering or interaction with MAPK-pathways. However, when the analysis of GR\textsuperscript{dim} mice was expanded to other inflammatory processes, we observed a failure of GCs to exert a full anti-inflammatory response. This was the case for exogenous actions of GCs in contact allergy and for the modulatory role of endogenous GCs in septic shock and for the early response during wound healing exhibiting elevated cytokine and chemokine expression (Grose et al., 2002; Tuckermann et al., 2007). As a side effect GC inhibition of bone formation occur in these mice. Keeping the findings in GR\textsuperscript{dim} mice in mind, what is the prediction for dissociating ligands, addressing the monomer function of the GR, the SEGRAs? Are they really potent to maintain therapeutic efficacy and reduce side effects? Although we do not know if the selective GR agonists (SEGRAs) do exactly mimick the GR\textsuperscript{dim} mutation on the molecular level one could speculate that their success – dissociating...
side effects from beneficial effects - depends on which conditions they are used for (Table 2). Whereas for contact allergy they may be less efficient, irritant dermatitis evoked by croton oil could be successfully cured as demonstrated (Schacke et al., 2004; Vayssiere et al., 1997) (Table 2). We would expect that effects on glucose homeostasis could be avoided with these compounds, but we would not assume that glucocorticoid induced osteoporosis would be prevented. However, some important inflammatory processes that are classically cured with steroids have not yet been analyzed with a SEGRA application or in the GR$^\text{dm}$ mice (Table 1 and 2). Also not all possible side effects have been extensively studied. Most importantly, the molecular mechanisms of GC suppression of asthma and rheumatoid arthritis have to be unraveled. Future research on these GC affected processes will tell if dissociating GR ligands are the solution for a low side-effect-therapy of these severe diseases affecting a major fraction of the population.

**References:**


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Figure Legends

Figure 1: Dimer dependent and independent control of gene expression by the GR. Schematic representation of the two major mechanisms for the regulation of gene expression by the ligand bound GR. On the left the cartoon shows the GR dimer bound to the palindromic GRE. This complex transactivates genes by interaction of the helix 12 of the GR with the LXXL motif present in the NR interaction box of co-activators of the p160 family. Interaction and recruitment of CBP/p300 coactivator/histone acetyl transferase (HAT) and numerous other coactivator complexes and HATs finally leads to decondensation of chromatin and enhanced transcriptional synthesis of mRNA molecules of the GR target gene. On the right the dimerization and DNA binding independent mechanism is shown. The GR molecule can interfere either with the activity of signal transduction components by direct or indirect mechanisms resulting in an inhibition of p38 and JNK activity or an enhancement of PI3K activity. In the nucleus the monomeric GR can influence the activity of pro-inflammatory transcription factors NF-κB, AP-1, IRF-3 and other factors. Co-integrators mediate this interaction as the nuclear isoform of the Trip-6 protein and STAMP or by competing out co-activators, such as the p160 family protein GRIP1. Traditionally interference of pro-inflammatory transcription factors is regarded as the molecular basis of anti-inflammatory effects of GC (Dr. Jekyll), whereas DNA binding contributes to the side effects of GC (Mr. Hyde). Recent studies show that for side effects such as glucocorticoid induced osteoporosis tethering might be sufficient and for full therapeutic efficacy of GC in contact dermatitis and for the endogenous actions of GC also dimerization induced DNA binding is required.
Figure 2: Contact hypersensitivity (CHS) responses are suppressed by GC via the GR in macrophages, require dimerization of the GR and the suppression of cytokines and chemokines. CHS is evoked by haptens, which are at least two times applied to the skin and therefore involve two phases: sensitization and elicitation. After the first exposure to the hapten “immature” dendritic cells of the dermis (DC) take up the hapten-protein complexes (1), and migrate to the draining lymph node (2). There the mature DC present the hapten by their surface MHC type II molecules to naïve T cells that recognize a matching hapten/MHCII complex with their T cell receptor and are sensitized by costimulation via B7 molecules from DCs by interaction of CD28 molecules (3). Sensitized T cells proliferate and repopulate the body (4). The elicitation phase starts by the second exposure of the same hapten to the skin (5). The endothelium (Endothel) becomes activated and allows the rapid entry of sensitized T cells (6), which in the skin are potently activated by hapten presenting DC (7). Those T cells trigger a massive activation of macrophages in the skin, which in turn release inflammatory mediators (8). These mediators (IL-1b, MCP-2, MIP-1 IP-10) lead to an influx of leukocytes, such as neutrophils (PMN) and monocytes (Mono) and manifest an inflammatory edema (9). The suppressive effect of GCs (10) is critical in macrophages and requires dimerization of the GR, since GR^lysMCre mice and GR^dim mice are resistant against GC suppression of CHS. GC action in keratinocytes and T cells is not sufficient, because mice lacking the GR in those cells (GR^ckCre, GR^K14Cre) can be cured with GCs.
### Phenotypes of pharmacological GC administration in GR<sup>dim</sup> mice

<table>
<thead>
<tr>
<th>Process</th>
<th>Phenotype</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Beneficial Effects of GC</strong></td>
<td></td>
</tr>
<tr>
<td>Treatment of irritant dermatitis</td>
<td>like in wild type (Reichardt et al., 2001)</td>
</tr>
<tr>
<td>Treatment of CHS</td>
<td>impaired (Tuckermann et al., 2007)</td>
</tr>
<tr>
<td>Suppression of inflammatory mediators</td>
<td>like in wild type: TNFa (Tuckermann et al., 2007), MMP-13, MMP-9 (Tuckermann et al., 1999) impaired: MCP-1, IP-10, IL-1b (Grose et al., 2002; Tuckermann et al., 2007)</td>
</tr>
<tr>
<td>PI3K coactivation by high dose GC, important for steroid protection from stroke</td>
<td>like in wild type: PI3K activity normal in GR&lt;sup&gt;dim&lt;/sup&gt; MEFs (Limbourg et al., 2002)</td>
</tr>
<tr>
<td><strong>Side Effects</strong></td>
<td></td>
</tr>
<tr>
<td>Induction of catabolic enzymes</td>
<td>impaired: TAT, PEPCK (Reichardt et al., 1998), (unpubl.)</td>
</tr>
<tr>
<td>GC induced osteoporosis</td>
<td>like in wild type (Tuckermann, unpubl.)</td>
</tr>
<tr>
<td><strong>Endogenous Actions of GC</strong></td>
<td></td>
</tr>
<tr>
<td>Survival of Sepsis</td>
<td>impaired (Kleyman et al., unpubl.)</td>
</tr>
<tr>
<td>Wound healing</td>
<td>like in wild type; but delayed kinetics (Grose et al., 2002)</td>
</tr>
<tr>
<td>Hepatic GH activity controlling body growth</td>
<td>like in wild type (Tronche et al., 2004)</td>
</tr>
</tbody>
</table>

Table 1: Phenotypes of pharmacological GC administration in GR<sup>dim</sup> mice
**Examples of beneficial and side effects of SEGRAs in vivo**

<table>
<thead>
<tr>
<th>SEGRA-Compound</th>
<th>Anti-inflammatory effect</th>
<th>Adverse effect</th>
</tr>
</thead>
<tbody>
<tr>
<td>RU24858 (Belvisi et al., 2001; Vayssiere et al., 1997)</td>
<td>croton-oil induced ear edema <em>efficiently reduced</em>, cotton-pellet granuloma model <em>efficiently reduced</em>, Seqhadox-model of lung edema <em>efficiently reduced</em></td>
<td>body weight loss <em>not changed</em>, thymus involution <em>not changed</em>, osteopenia <em>not changed</em></td>
</tr>
<tr>
<td>ZK216348 (Schacke et al., 2004)</td>
<td>croton-oil induced ear edema <em>efficiently reduced</em></td>
<td>body weight reduction <em>decreased</em>, blood glucose elevation <em>decreased</em>, spleen involution <em>decreased</em>, skin atrophy <em>slightly decreased</em>, adrenocorticotropic hormone suppression <em>not changed</em></td>
</tr>
<tr>
<td>AL-438 (Coghan et al., 2003)</td>
<td>carrageenan-induced arthritis, adjuvant induced arthritis <em>efficiently reduced</em></td>
<td>hyperglycemia <em>decreased</em>, inhibition of bone apposition <em>decreased</em></td>
</tr>
</tbody>
</table>

Table 2: Examples of SEGRAs tested in vivo for anti-inflammatory activities and adverse effects in comparisons with conventional GCs
Acknowledgements:

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Dimerization-induced DNA Binding

Dimerization-independent Gene regulation

Nucleus

p160
LXXLL
Pol
GRE
Pol
AP-1
NF-kB
IRF-3

Dimerization-induced DNA Binding
mRNA

Dimerization-independent Gene regulation

Nucleus

p38
JNK
PI3K

p38
JNK
PI3K

Side effects?
Mr. Hyde

Therapy?
Dr. Jekyll

glucocorticoid induced osteoporosis

Contact Allergy
Septic shock

Fig. 1
Fig. 2

1. Keratinocytes
2. Endoth
3. Lymphnode
4. DC
5. T
6. T
7. T
8. IL-1β
9. IP-10
10. MIP-2
11. MCP-1

Mφ
GCs

Sensitization        Elicitation