Supplemental Experimental Procedures

Fly stocks

The line $P\{w^+, EP\}GE50159$ was obtained from GenExel, Korea. Deficiencies in heterochromatin 3 and the $\alpha4\text{-tub-Gal4}$ driver line are available from the Bloomington stock center. Transgenic lines with $His2Av\text{-mRFP1}$, $EGFP\text{-Cid}$ (Schuh et al., 2007), $gSMC3\text{-HA}$ (Heidmann et al., 2004), $201Y\text{-Gal4}$ (Yang et al., 1995), $H24\text{-Gal4}$ (Zars et al., 2000) and $Cha\text{-Gal4}$ (Salvaterra and Kitamoto, 2001) have been described previously. A $mhc\text{-Gal80}$ fly stock (C. Winter and L. Luo, unpublished) was kindly provided by Liqun Luo. A complete stocklist with all genotypes and abbreviations used in this paper can be found in Supplemental Table S1.

Generation of Rad21 alleles

The insertion site of the homozygous viable $P\{w^+, EP\}GE50159$ line 4 kb upstream of the transcription start site of $Rad21$ was confirmed by inverse PCR according to a standard protocol. Imprecise excisions were generated by crossing the GE50159 line to flies expressing a stable source of the P-element transposase. Out of 500 excision events, 23 homozygous lethal lines were isolated. 4 independently generated deletions affecting $Rad21$ were subsequently identified by PCR ($Rad21^{ex3}$, $Rad21^{ex8}$, $Rad21^{ex15}$, $Rad21^{ex16}$) and confirmed by sequencing DNA fragments spanning the breakpoints.
**Generation of transgenic flies expressing TEV protease**

To generate a NLS- and v5-epitope-tagged TEV protease expression construct (nuclear TEV protease), one N-terminal consensus sequence of the SV40 nuclear localization signal (NLS) followed by one v5-epitope tag were introduced by PCR at the 5’end of the coding sequence of TEV-NLS2, using the yeast vector YIplac204 (Uhlmann et al., 2000) as template. Primer sequences are listed below. The PCR product NLS-v5-TEV-NLS2 was cloned as EcoRI/NotI fragment into the pUAST (Brand and Perrimon, 1993) or pCaSpeR-hs (Thummel, 1992) transformation vectors to obtain UAS-NLS-v5-TEV-NLS2 or hs-NLS-v5-TEV-NLS2, respectively. Transgenic lines were produced by standard P-element-mediated germline transformation (Rubin and Spradling, 1983) and either recombined to the Rad21ex3 allele (Rad21ex3, UAS-TEV/TM3, Sb, Kr-Gal4, UAS-GFP) or crossed into the Rad21ex3 background (hs-TEV; Rad21ex3/TM6B, Tb).

**Generation of flies surviving on transgenic Rad21^{TEV}**

C-terminally 10xmyc-tagged Rad21 was created based on the EST clone LD14219 obtained from the Berkeley Drosophila Genome Project (BDGP). The coding sequence of the 10xmyc-tag was amplified by PCR from the plasmid gthr-myc (Leismann et al., 2000) and cloned as Bst-BI fragment into the unique Bst-BI site in LD14219, located 12 nucleotides upstream of the Rad21 translational stop codon. The sequence encompassing Rad21-myc10 was excised as a 2790 bp Eco-RI//Kpn-I fragment and cloned into pUAST (Brand and Perrimon, 1993) to obtain pUAS-Rad21-\textit{myc}_{10}. 
To generate TEV-cleavable versions of Rad21 (Rad21\textsuperscript{TEV}), SpeI-restriction sites were introduced into the coding region of \textit{Rad21} after amino acids 175, 197, 271 or 550 by site-directed PCR-mutagenesis, using \textit{pUAS-Rad21-myc}\textsubscript{10} as template (primer sequences see below). An AvrII/NheI restriction fragment encoding 3 tandem arrays of the TEV-recognition sequence ENLYFQG (kindly provided by Stephan Gruber, for details on the sequence see below) was inserted into the newly generated SpeI-site. The \textit{Rad21(3TEV)-myc\textsubscript{10}} fragment (EcoRI/KpnI-blunt) was introduced into the multiple cloning site (MCS) of a modified EcoRI/blunt cut pBS vector containing a MCS and 3’UTR flanked by FRT-sites (<). Next, the sequence comprising \textless Rad21(3TEV)-myc\textsubscript{10}-3’UTR \textgreater{} was excised with KpnI and inserted into a KpnI-cut modified pCaSpeR transformation vector (derived from 10xUAS, G. Dietzl), in which the 10xUAS-sequence had been replaced with the sequence of the tubulin-promotor (derived from plasmid M>P\textsuperscript{2}, Casali and Struhl, 2004) (final vectors: tubpr<Rad21(3TEV)-myc\textsubscript{10}-3’UTR<SV40).

To generate vectors expressing myc-tagged Rad21 without TEV-cleavage sites, the open reading frame of \textit{Rad21-myc\textsubscript{10}} was PCR-amplified from \textit{pUAS-Rad21-myc\textsubscript{10}} and introduced into the same modified vector (final vector: tubpr-Rad21-myc\textsubscript{10}-SV40).

Transgenic lines were produced by standard P-element-mediated germline transformation (Rubin and Spradling, 1983). Transgenes were tested for their ability to rescue the lethality of \textit{Rad21} null mutations. Transgenes with TEV-cleavage sites at positions 271 or 550 and a transgene without TEV-sites were functional.

\textbf{Immunoblotting}
For the preparation of embryonic extracts, dechorionated embryos were homogenized in SDS-sample loading buffer 3-6 hours after egg deposition. Proteins were resolved by SDS-polyacrylamide gel electrophoresis and transferred to a PVDF membrane. The blot was probed with mouse anti-myc 9E10 (1:15, Sigma-Aldrich) and mouse anti-α-tubulin (DM1A) (1:20000, Sigma-Aldrich) using ECL detection (Amersham). Protein extracts from pupae, 3rd instar larval salivary glands and 3rd instar larvae without salivary glands were prepared after dissection in PBS essentially as described in Experimental Procedures.

**Immunostaining of polytene chromosome squashes**

Salivary glands from wandering 3rd instar larvae were dissected in PBS, permeabilized in PBX* (PBS, 0.1% Triton X-100, 3.7% formaldehyde) for 30 seconds, fixed in 45% acetic acid/3.7% formaldehyde for 5-7 minutes and squashed according to standard procedures. Slides were blocked for 1 hour at room temperature in PBST + 5% BSA (PBS, 0.01% Tween20, 5% BSA) and incubated with primary antibodies (diluted in blocking solution) overnight at 4°C. After washing in PBST (3x10 minutes), slides were incubated with Alexa-conjugated secondary antibodies at room temperature for 1.5 hours, washed as before and mounted using VECTASHIELD mounting medium containing DAPI (Vector Laboratories).

The following (additional) primary antibodies were used: guinea-pig anti-SA (1:100, Dorsett et al., 2005), rabbit anti-SMC1 (1:100, Dorsett et al., 2005), rat anti-SMC1 (1:500, Malmanche et al., 2007), mouse anti-HA 16B12 (1:250, Covance), mouse anti-polymerase II 8WG16 (1:20, Covance), rabbit anti-trx (1:50, Chinwalla et al., 1995), rabbit anti-Pc (1:200, Zink and Paro, 1989), rabbit anti-HSF (1:80, Andrilis et al., 2000), rabbit anti-BEAF32 (SCBP) (1:50, Zhao et al., 1995) and mouse anti-Z4
Primer and DNA sequences

Restriction enzyme sites are shown in lower case, the start codon ATG in bold, the v5-epitope sequence in italics, the NLS sequence underlined.

**NLS-v5-TEV-NLS**

**AP2f**

5′-AAAgaattcAAAATGCTAAGAAAAAGGAGGTTGCATCCGTAAGCCTATCCCTAACCTCTCTCTCGGTCTCGATTCTACGGGAGAAAGCTTTGAAGGACCAGC-3′

**AP4r**

5′-AAAAACCCTCGAGGCCcgcccgccAG-3′

**pUAS-Rad21-myc**

**SH28**

5′-CGATTATTCGAAAAACCCAAATTGTTGCATGGATG

G-3′

**SH29**

5′-CGATTA TTGAATGCCCATGTTCGCCCAAG-3′

**tubpr-Rad21-myc** and **tubpr-Rad21(3TEV)-myc-3′ UTR**

**AP28-EcoRIRad21f**

5′-AAAgaattcAAAATGTTCTATGAGCAGCATTATTTTCAG-3′

**AP29-Rad21NotIr**

5′-AAAgcggccgcaATTAACACAGATTTACATTTCAAC-3′

**AP20-TEV175SpeIf**

5′-CCTCTATTTTCAAGGAGTCTCTGATTCAATGGGACACTATCTCCTGCTCTATATTTGACGATATA-3′

**AP21-175NheISpeIr**

5′-actagtGCCCTTGGAAAAATAGAGCTCGGTCTGCTCTGTTAATTCGATTG-3′

**AP22-197SpeIf**

5′-TTTTGGAAAATATTGAGACTAGTCTCTGAGTCCCATGAGC-3′

**AP23-197SpeIr**

5′-ACTAGCTGCTTACGACTGTCATGAGC-3′

**AP24-271SpeIf**

5′-CATAATGTCCCCTCCTCACTAGCAGGAGACTCACATGAGC-3′

**AP25-271SpeIr**

5′-ACTAGCTGCTTACGACTGTCATGAGC-3′

**AP26-550SpeIf**

5′-ACTAGCTGCTTACGACTGTCATGAGC-3′

**AP27-550SpeIr**

5′-ACTAGCTGCTTACGACTGTCATGAGC-3′

3 TEV recognition sites (3TEV)
Supplemental References

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## Table S1: Strains used in this study

| Genotype | Abbreviation | Source |
|----------|--------------|--------|
| w<sup>1118</sup> | w | Bloomington stock centre |
| w; lethal/CyO; TM2, {∆2-3}/Sb, {∆2-3} | P-element transposase | kindly provided by Frank Schnorrer |
| w; P[w+, EP]GE50159 (III) | GE50159 | GenExel, Korea |

### Gal4 driver

| Genotype | Abbreviation | Source |
|----------|--------------|--------|
| w; act-Gal4/CyO | act-Gal4 | Barry Dickson lab stocks |
| w; hs-Gal4 (III) | hs-Gal4 | Barry Dickson lab stocks |
| w; P[w+, mata4-tub-Gal4-VP16] | a4-tub-Gal4 | Bloomington stock centre |
| w*; P[w+, G4B]F4 (II) | F4-Gal4 | Weiss et al., 1998 |
| w; elav-Gal4 (3AF) (III) | elav-Gal4 | Barry Dickson lab stocks |
| w; nSyb-Gal4/CyO | nSyb-Gal4 | kindly provided by Julie Simpson |
| y, w; 201Y-Gal4, UAS-mCD8-GFP/CyO; MKRS, Sb/TM6B, Tb | 201Y-Gal4, UAS-mCD8-GFP | Liqun Luo lab stocks |
| w; Pin/CyO; H24-Gal4, UAS-mCD8-GFP | H24-Gal4, UAS-mCD8-GFP | Liqun Luo lab stocks |
| w; Cha-Gal4 (II) | Cha-Gal4 | Salvaterra and Kitamoto, 2001 |

### Transgenes and Mutants

| Genotype | Abbreviation | Source |
|----------|--------------|--------|
| w*; P[w+, His2Av-mRFP1] (II.2) | His2Av-mRFP1 | Schuh et al., 2007 |
| w*; P[w+, EGFP-Cid] (II.1) | EGFP-Cid | Schuh et al., 2007 |
| w*; P[w+, gSMC3-HA12] (III.2) | gSMC3-HA | Heidmann et al., 2004 |
| y, w; P[w+, mhc-Gal80] (III) | mhc-Gal80 | C. Winter and L. Luo, unpublished |
| Df(3L)2-66, kn[ri-1] p[p]/TM3, Sb, Ser | Df(3L)2-66 | Bloomington stock centre |

### Rad21-excisions

| Genotype | Abbreviation | Source |
|----------|--------------|--------|
| w; Rad21<sup>ex3</sup>/TM3, Sb, Kr-Gal4, UAS-GFP | Rad21<sup>ex3</sup> | present study |
| w; Rad21<sup>ex8</sup>/TM3, Sb, Kr-Gal4, UAS-GFP | Rad21<sup>ex8</sup> | present study |
| y, w; Rad21<sup>ex15</sup>/TM3, Sb, Kr-Gal4, UAS-GFP | Rad21<sup>ex15</sup> | present study |
| w; Rad21<sup>ex16</sup>/TM3, Sb, Kr-Gal4, UAS-GFP | Rad21<sup>ex16</sup> | present study |
### TEV protease transgenes

| w; P(w+, UAS-NLS-v5-TEV-NLS2) (III) | UAS-TEV | present study |
| w; P(w+, hs-NLS-v5-TEV-NLS2) (II) | hs-TEV | present study |

### Rad21-excision + TEV-protease

| w; Rad21ex3, P(w+, UAS-NLS-v5-TEV-NLS2)/TM3, Sb, Kr-Gal4, UAS-GFP | Rad21ex3, UAS-TEV/TM3, Sb, Kr>GFP | present study |
| w; Rad21ex3, P(w+, UAS-NLS-v5-TEV-NLS2), hs-Gal4/TM3, Sb, Kr-Gal4, UAS-GFP | Rad21ex3, UAS-TEV, hs-Gal4/TM3, Sb, Kr>GFP | present study |
| w; P(w+, hs-NLS-v5-TEV-NLS2); Rad21ex3/TM6B, Tbx, ubiquitin-GFP | hs-TEV; Rad21ex3/TM6B, Tbx | present study |

### transgenic Rad21 (+/- TEV-sites)

| w; P(w+, tubpr<Rad21(550-3TEV)-myc10<SV40> (III) | Rad21(550-3TEV)-myc (Rad21TEV) | present study |
| w; P(w+, tubpr<Rad21(271-3TEV)-myc10<SV40> (III) | Rad21(271-3TEV)-myc (Rad21TEV) | present study |
| w; P(w+, tubpr-Rad21-myc10<SV40> (III) | Rad21-myc (Rad21) | present study |
| w; P(w+, tubpr<Rad21(271-3TEV)-myc10<SV40> (II.3) | Rad21(271-3TEV)-myc (Rad21TEV) | present study |
| w; P(w+, tubpr<Rad21(271-3TEV)-myc10<SV40> (II.7) | Rad21(271-3TEV)-myc (Rad21TEV) | present study |
| w*, P(w+, tubpr<Rad21(271-3TEV)-myc10<SV40> (II.3), P(w+, tubpr<Rad21(271-3TEV)-myc10<SV40> (II.7) 2x Rad21(271-3TEV)-myc (2x Rad21TEV) | present study |

### Rad21-excision + transgenic Rad21 (+/- TEV-sites)

| w; Rad21ex15, P(w+, tubpr<Rad21(550-3TEV)-myc10<SV40> (III) | Rad21ex15, Rad21TEV | present study |
| w; Rad21ex8, P(w+, tubpr<Rad21(271-3TEV)-myc10<SV40> (III) | Rad21ex8, Rad21TEV | present study |
| w; P(w+, tubpr<Rad21(271-3TEV)-myc10<SV40> (III), P(w+, tubpr<Rad21(271-3TEV)-myc10<SV40> (II.3), P(w+, tubpr<Rad21(271-3TEV)-myc10<SV40> (II.7); Rad21ex3 2x Rad21(271-3TEV)-myc (2x Rad21TEV) | present study |

### Rad21-excision + Gal4 (+ TEV-protease or transgenic Rad21TEV)

| w; Rad21ex3, hs-Gal4/TM3, Sb, Kr-Gal4, UAS-GFP | Rad21ex3, hs-Gal4/TM3, Kr-Gal4, UAS-GFP | present study |
| w*; P(w+, α4-tub-Gal4-VP16)/CyO, wg-lacZ; Rad21ex3/TM3, Sb, ubx-lacZ | α4-tub-Gal4/CyO, wg-lacZ; Rad21ex3/TM3, ubx-lacZ | present study |
| w*; P(w+, α4-tub-Gal4-VP16), P(w+, His2Av-mRFP1) (II.2), P(w+, EGFP-Cid) (II.1)/CyO, wg-lacZ; Rad21ex3/TM3, Sb, ubx-lacZ | α4-tub-Gal4, His2Av-mRFP1, EGFP-Cid/CyO; Rad21ex3/TM3, ubx-lacZ | present study |
| w; P[w+, GawB]F4/CyO, Kr-Gal4, UAS-GFP; Rad21\textsuperscript{ex15}, P[w\textsuperscript{+}], tubpr<Rad21(550-3TEV)-myc\textsubscript{10}<SV40) | F4-Gal4/CyO, Kr-Gal4, UAS-GFP; Rad21\textsuperscript{ex15}, Rad21(550-3TEV)-myc | present study |
| w; Rad21\textsuperscript{ex3}, P[w\textsuperscript{+}, UAS-NLS-v5-TEV-NLS\textsubscript{10}], elav-Gal4/TM3, Sb, Kr-Gal4, UAS-GFP | Rad21\textsuperscript{ex3}, UAS-TEV, elav-Gal/VM3, Kr-Gal4, UAS-GFP | present study |
| w; nsyb-Gal4/CyO; Rad21\textsuperscript{ex3}, P[w\textsuperscript{+}, UAS-NLS-v5-TEV-NLS\textsubscript{10}]/TM3, Sb, Kr-Gal4, UAS-GFP | nsyb-Gal4/CyO; Rad21\textsuperscript{ex3}, UAS-TEV/TM3, Kr-Gal4, UAS-GFP | present study |
| w; Rad21\textsuperscript{ex3}, P[w\textsuperscript{+}, UAS-NLS-v5-TEV-NLS\textsubscript{10}], H24-Gal4, UAS-mCD8-GFP/TM6B, Tb | Rad21\textsuperscript{ex3}, UAS-TEV, H24-Gal4, UAS-mCD8-GFP/TM6B, Tb | present study |
| w; 201Y-Gal4, UAS-mCD8-GFP/(CyO); Rad21\textsuperscript{ex3}, P[w\textsuperscript{+}, tubpr<Rad21(550-3TEV)-myc\textsubscript{10}<SV40) | 201Y-Gal4, UAS-mCD8-GFP; Rad21\textsuperscript{ex3}, Rad21 | present study |
| w; 201Y-Gal4, UAS-mCD8-GFP/(CyO); Rad21\textsuperscript{ex3}, P[w\textsuperscript{+}, tubpr-Rad21(550-3TEV)-myc\textsubscript{10}<SV40) | 201Y-Gal4, UAS-mCD8-GFP; Rad21\textsuperscript{ex3}, Rad21 | present study |
| w; 201Y-Gal4, UAS-mCD8-GFP/CyO; Rad21\textsuperscript{ex3}, P[w\textsuperscript{+}, mhc-Gal80], P[w\textsuperscript{+}, UAS-NLS-v5-TEV-NLS\textsubscript{10}]/TM6B, Tb | 201Y-Gal4, UAS-mCD8-GFP/CyO; Rad21\textsuperscript{ex3}, mhc-Gal80, UAS-TEV/TM6B, Tb | present study |
| w; Cha-Gal4; Rad21\textsuperscript{ex3}, P[w\textsuperscript{+}, UAS-NLS-v5-TEV-NLS\textsubscript{10}]/TM6B, Tb | Cha-Gal4; Rad21\textsuperscript{ex3}, UAS-TEV/TM6B, Tb | present study |
| w; UAS-CBD8-GFP; If/CyO; Rad21\textsuperscript{ex15}, P[w\textsuperscript{+}, tubpr<Rad21(550-3TEV)-myc\textsubscript{10}<SV40) | UAS-CBD8-GFP; If/CyO; Rad21\textsuperscript{ex15}, Rad21 | present study |
| w*; Rad21\textsuperscript{ex3}, P[w\textsuperscript{+}, tubpr-Rad21(550-3TEV)-myc\textsubscript{10}<SV40) | Rad21\textsuperscript{ex3}, Rad21, UAS-TEV/TM6B, Tb | present study |

**Table S2: Rescue of Rad21 excision alleles by ectopic expression of either Rad21-myc or Rad21\textsuperscript{TEV}-myc**

| Relative Viability * | Rad21\textsuperscript{ex16}/TM3 Sb, Kr>GFP | Rad21\textsuperscript{ex3}/TM3 Sb, Kr>GFP |
|---------------------|---------------------------------|---------------------------------|
| Pupae#              | Adults##                        | Pupae#                          | Adults##                        |
| ♀ Rad21\textsuperscript{ex16}/TM3 Sb, Kr>GFP | 0                               | 0                               | 0                               |
| ♀ Rad21\textsuperscript{ex3}/TM3 Sb, Kr>GFP | 0                               | 0                               | 0                               |
| ♀ Rad21\textsuperscript{ex3}, Rad21-myc   | 95.1                            | 101                             | 105                             | 93.9                             |
| ♀ Rad21\textsuperscript{ex15}, Rad21(550-3TEV)-myc | 111                             | 81.4                            | 107                             | 78.4                             |
| ♀ Rad21\textsuperscript{ex8}, Rad21(271-3TEV)-myc | 87.5                            | 96.6                            | 90.6                            | 96.4                             |

* Relative Viability: percentage of rescued pupae/adults, normalized to the values obtained when Rad21\textsuperscript{ex}/TM3 Sb, Kr>GFP males were crossed to w\textsuperscript{1118} females.

# Rescued pupae were identified by the absence of GFP-expression (n ≥ 400).

## Rescued adults were identified by the absence of Sb (n ≥ 250).
Supplemental Figures

Figure S1. Multiple sequence alignment of protein-sequences of metazoan Rad21-homologs.

Annotated Rad21-protein sequences or sequence fragments were aligned to each other using ClustalW (Alexander Schleiffer, unpublished data). The conserved N- and C-terminal domains (green frames), human and (predicted) D. melanogaster separase cleavage sites (blue and red asterisks, respectively) and the boundaries of exons 2-3 and 3-4 are indicated. Four poorly conserved regions were chosen to introduce 3 tandem arrays of TEV recognition sequences (black arrows). The amino acid position after which the TEV sites had been introduced is indicated. TEV sites that rendered a functional Rad21\textsuperscript{TEV} protein (271 and 550) are highlighted in purple boxes.
Figure S2. Rad21<sup>TEV</sup> is cleaved by zygotically expressed TEV before mitosis.

(A) Total extracts corresponding to 30 or 10 embryos were analyzed by immunoblotting with antibodies against myc and α-tubulin 3-6 hours after egg deposition. (-TEV) Embryos surviving on myc-tagged Rad21<sup>TEV</sup> express maternal Gal4, but do not contain the UAS-TEV transgene. (+TEV) Embryos surviving on myc-tagged Rad21<sup>TEV</sup> express maternal Gal4, which drives zygotic TEV
expression. The arrowhead indicates the position of full-length myc-tagged Rad21\textsuperscript{TEV}, the asterisk that of the C-terminal TEV-cleavage product. Note that only 50% of the embryos used for the +TEV extracts contain the UAS-TEV construct (for details on genetic crosses see Supplemental Experimental Procedures). Therefore, the data is fully consistent with complete or almost complete cleavage of Rad21\textsuperscript{TEV}.

(B) Unstable kinetochore attachment after TEV expression. Frames shown were taken at 15 second intervals from a cell during the mitotic arrest resulting from zygotic TEV expression in embryos surviving on Rad21\textsuperscript{TEV}. DNA is marked with His2Av-mRFP1 (red), kinetochores with EGFP-Cid (green). The arrow follows the movement of a single kinetochore.

(C) Frames were taken at times indicated (min:sec) from a cell in a TEV-expressing embryo surviving on Rad21\textsuperscript{TEV}. DNA is marked with His2Av-mRFP1 (red), kinetochores with EGFP-Cid (green). After an initial mitotic arrest, chromosome decondensation starts abruptly (16:00) and unattached chromatids in the central region are cut by the cleavage furrow.
Figure S3. The cohesin complex binds to polytene chromosomes.

(A) Polytene chromosomes from wild-type flies (top two rows) or flies transgenic for HA-tagged SMC3 (bottom row) were coimmunostained with antibodies against endogenous Rad21 (green) and either endogenous SA (top row), endogenous SMC1 (middle row) or the HA-epitope (bottom row) (red). A schematic of the cohesin complex is shown at the left of each row (arrow points to the subunit costained with DAPI).
Rad21). DNA was visualized with DAPI. Scale bars, 15µm.

(B) Polytenic chromosome spreads from 3rd instar larvae, which express heat-inducible TEV (hs-TEV) and myc-tagged Rad21\textsuperscript{TEV} as their only source of Rad21, were prepared after a 45 min heat shock. Spreads were coimmunostained with antibodies against endogenous Rad21 and myc (top) or myc (green) and SMC1 (red) (bottom). Note the absence of Rad21-, myc- and SMC1-staining after TEV cleavage of Rad21\textsuperscript{TEV}. DNA was visualized with DAPI. Scale bar, 15µm (top), 10µm (bottom).
Figure S4. Rad21 binds to distinct regions on polytene chromosomes.

(A) Polytene chromosomes from wild-type flies (w^{1118}) were coimmunostained with...
antibodies against Rad21 (green) and BEAF (red), a well-characterized boundary-associated factor. DNA is shown in blue in merged images. Two different magnifications are shown (top row: 40x objective; bottom row: 100x objective, split channels). BEAF and Rad21 localize to distinct interband regions. Scale bar, 20µm.

(B) Polytene chromosomes from wild-type flies (w^{1118}) were coimmunostained with antibodies against Rad21 and (a) PolII (RNA Polymerase II), (b) HSF (Heat-Shock Factor), (c) Z4 (interband-specific Zinc-finger protein), (d) Pc (Polycomb) and (e) trx (Trithorax). DNA was visualized with DAPI. As seen in the merged images (Rad21 in green, other proteins in red), the distribution of cohesin and of the other tested factors differs significantly. White arrows point to the few regions, in which an overlap between Rad21 and the other factor tested could be detected. The insets in a1-a4 show a higher magnification (3x) of a PolII-stained chromosomal puff, which is flanked by Rad21-bound regions. Scale bars, 20µm (a and b), 10µm (c-e).
Figure S5. Cohesin is required for salivary gland development.

(A) Salivary glands were dissected from 3rd instar larvae that had been raised at 18°C throughout development. (Left) In the whole-mount preparations, DNA was stained with DAPI. (Right) Polytene chromosome spreads were immunostained with antibodies against Rad21 (green). DNA was visualized with DAPI (blue). Three different genotypes were compared: a) gRad21/Rad21<sup>ex3</sup>, hs-Gal4, UAS-TEV; b) Rad21<sup>ex15</sup>, Rad21<sup>TEV</sup>/Rad21<sup>ex3</sup>, hs-Gal4; c) Rad21<sup>ex15</sup>, Rad21<sup>TEV</sup>/Rad21<sup>ex3</sup>, hs-Gal4, UAS-TEV. Since polytene chromosome spreads from genotypes (a) and (b) are similar, only a representative spread from (a) is shown. Note that salivary glands with reduced amounts of Rad21 have smaller but not fewer cells. Scale bar, 500µm (whole
mount salivary glands), 20µm (polytene chromosome spreads).

(B) Quantitative analysis of salivary glands average length (in mm), total number of cells per salivary gland and the average diameter per nucleus (in µm) was performed from salivary glands from 3rd instar larvae of the indicated genotypes (>10 per genotype). Larvae in which Rad21^{TEV} has been cleaved show smaller salivary glands with smaller nuclei. The number of cells per salivary glands remains unaltered.

(C) Protein extracts from a strain carrying hs-Gal4, UAS-TEV and surviving on myc-tagged Rad21^{TEV} were analyzed by Western Blotting. Extracts of larvae lacking salivary glands (L), dissected salivary glands (SG) or pupae (P) were prepared from crosses raised at 18°C. Samples from lanes 3 and 4 were prepared 1 hour after heat shock treatment (45 min 37°C). Western Blot analysis was performed with antibodies against myc (detecting full-length Rad21^{TEV}-myc (arrow) and the C-terminal TEV-cleavage fragment (arrowhead)), v5 (detecting TEV-protease) and actin (loading control). Before heat shock induction of TEV, significant levels of the protease could be detected in salivary glands (open circle). The TEV cleavage fragment of Rad21 is also observed. Neither TEV protease nor Rad21^{TEV} cleavage fragments were detected in larvae without salivary glands or pupae before TEV induction. A Molecular Weight Marker (in kDa) is shown on the left.
Figure S6. Analysis of pruning in γ neurons with H24-Gal4-induced TEV cleavage of Rad21.

H24-Gal4 was used to drive expression of TEV and mCD8 in γ neurons of the...
mushroom body from flies that survived on transgenic Rad21 with (Rad21^{TEV}) or without (Rad21) TEV cleavage sites. Shown are maximal Z projections of single confocal sections of a right brain hemisphere, stained with antibodies against mCD8 (green) and FasII (red). Scale bars, 20μm.

(A) Larval γ neurons from a Rad21^{TEV} brain project into the dorsal and medial lobes (filled green arrowheads).

(B) (24h APF, left) In the presence of Rad21 (control), γ neurons have pruned their medial and dorsal axon projections (open arrowheads) as well as their dendrites. In γ neurons of Rad21^{TEV} pupae, larval axon-projections persist in the medial and dorsal lobes (filled green arrowheads). Note also the presence of unpruned dendrites (green arrow). (4d APF, right) In the presence of Rad21 (control), γ neurons have re-extended their axons medially towards the midline. Axons of α/β neurons in the dorsal and medial lobes are labeled with FasII (red arrows). In γ neurons of Rad21^{TEV} pupae, larval axon-projections as well as dendrites (filled green arrowheads and green arrow, respectively) persist in the dorsal and medial lobe. Projections of α/β neurons are normal (red arrows).

(C) In the presence of Rad21 (top), γ neurons are tightly bundled and project exclusively towards the midline at 4d APF. No γ neurons are found in the FasII-positive dorsal lobe (open green arrowhead). In the absence of Rad21 (bottom), γ neuronal projections persist in the dorsal and medial lobes (filled green arrowheads), but are often disorganized and mistargeted. Although the FasII staining for α/β neurons appears weaker than in the control, the projection pattern is normal.
Figure S7. TEV cleavage of Rad21 in cholinergic-neurons does not cause mitotic defects.

Brains from control (-TEV: *UAS-CD8-GFP; Cha-Gal4/CyO; Rad21ex15, Rad21TEV/TM6B) and TEV-cleaved brains (+TEV: *UAS-CD8-GFP; Cha-Gal4/CyO; Rad21ex15, Rad21TEV/Rad21ex3, UAS-TEV) were immunostained for CD8 (green) and the mitotic marker phospho-histone H3 (PH3, red).

A) High magnification images show that there is no co-localization between the CD8 positive cells and the mitotic marker. Additionally, mitotic figures in TEV-cleaved brains look similar to the controls. Scale bars, 20µm.
B) Confocal images of brains and ventral nerve cords from both control and TEV-cleaved brains. PH3 staining reveals that there is no detectable accumulation of mitotic figures after TEV cleavage, and that there are no obvious morphological defects in the CD8 positive cholinergic neurons. Scale bars, 100µm.