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17. The Lebendun Nappe here contains three horizons of garnet amphibolite (8 to 15 m thick) interlayered with pelite. The "pelite" grades from a quartz-rich (86% SiO_2) psammite at the contact to a more typical pelite (75% SiO_2) within ~10 cm. The amphibolite contains biotite, which decreases in mode away from the contact, and garnet porphy-

roblasts up to 5 mm in diameter. At the contact, garnets are more sparse and rarely exceed 500 μm in diameter. Pelitic garnets continue to decrease in diameter and mode away from the contact, <200 μm near the contact and reaching zero after several meters.

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Spatial Patterns in the Distribution of Tropical Tree Species

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Fully mapped tree census plots of large area, 25 to 52 hectares, have now been completed at six different sites in tropical forests, including dry deciduous to wet evergreen forest on two continents. One of the main goals of these plots has been to evaluate spatial patterns in tropical tree populations. Here the degree of aggregation in the distribution of 1768 tree species is examined based on the average density of conspecific trees in circular neighborhoods around each tree. When all individuals larger than 1 centimeter in stem diameter were included, nearly every species was more aggregated than a random distribution. Considering only larger trees (≥ 10 centimeters in diameter), the pattern persisted, with most species being more aggregated than random. Rare species were more aggregated than common species. All six forests were very similar in all the particulars of these results.

The spatial dispersion of individuals in a species is central in ecological theory (1, 2). Patchiness, or the degree to which individuals are aggregated or dispersed, is crucial to how a species uses resources, to how it is used as a resource, and to its reproductive biology. Spatial patterns have been a particularly important theme in tropical ecology, because high diversity in the tropics begets low densities. Since Wallace (3) noted how difficult it was to find two individuals of the same species, the hyperdispersion of tropical trees has focused much of theoretical tropical ecology.

In 1979, Hubbell (4) published a large study of dispersion of trees in a dry forest in Costa Rica. His results were contrary to Wallace's long-prevailing wisdom and the Janzen-Connell prediction (5, 6) that wide dispersion is a defense against predators. Most species were aggregated, so that near neighborhoods of a tree had a higher than average density of conspecifics. Since that study,

though, contradictory results have appeared, particularly from Lieberman and Lieberman (7), who found that most species in a wet forest in Costa Rica, as well as from a literature survey, were not aggregated.

Over the past two decades, we have been assembling a long-term, large-scale, global research effort on spatial patterns and dynamics of tropical forests (8, 9). An international team has now fully censused six plots in five tropical countries, mapping and identifying every individual of ≥ 1 cm in stem diameter over 25 to 52 ha at each plot (Table 1). The large plot size is necessary to encompass substantial populations of most tree species in the community. Major goals of this effort have been to examine Janzen-Connell effects, density dependence, and the spacing pattern of individual species.

The six sites represent a wide variety of tropical forests (Table 1). At one extreme, the two plots in Malaysia are in tall, evergreen forest; have no regular dry season; and include

over 800 tree and treelet species each. The Sinharaja forest is also very wet and evergreen, but its island setting reduces species diversity. The site in India is in dry forest with a fairly open canopy, grassy understory, and just 70 species; the Thai site is also dry and low in diversity. The single site in Central America is moist forest, structurally quite like the Malaysian sites, but intermediate in climate and diversity. The forests also cover a wide taxonomic range. Four of the Asian sites are dominated by the family Dipterocarpaceae, but few species are shared among them. The Indian and American sites are distinct taxonomically; they are not dominated by a single family and have few (India) or no (Panama) dipterocarps.

We evaluate spatial patterning by examining neighborhoods around individual trees. For each individual, we tallied the number of conspecifics between x and $x + \Delta x$ meters for all $x + \Delta x$ inside the plot. We also calculated the area inside the plot of each of these annuli. The number of neighbors N_x and the area A_x in each annulus at distance x were then summed over all individuals of a given species. $D_x = \Sigma N_x / \Sigma A_x$ gives the density of neighboring conspecifics as a function of distance from the average individual. This is a biologically meaningful measure of clumping, because it evaluates the conspecific population density in the neighborhood of

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each tree. It is closely related to Ripley's K statistic [called the correlation integral by astronomers when applied, in three dimensions, to the distribution of galaxies (10)], but K is a cumulative distribution, whereas our neighborhood density is a probability density function; that is, K_x refers to conspecifics of $<x$ meters from the focal tree, and D_x refers to an annulus between x and $x + \Delta x$ meters. Although the K statistic is very popular (11), our approach has the advantage of isolating specific distance classes, whereas K confounds effects at larger distances with effects at shorter distances (12, 13).

To compare species with various population densities, we standardized D_x by dividing it by the mean density of a given species across the whole plot. We call this standardized index the relative neighborhood density, or Ω . In a perfectly random distribution, $\Omega_x = 1$ for all distances x . Aggregation is indicated when $\Omega_x > 1$ at short distances, whereas $\Omega_x < 1$ at short distances indicates spacing at some scale, or hyperdispersion. A great advantage of this standardized statistic is that it is sample-size independent, which allowed us to directly compare species and stem diameter classes and offered a bootstrap method for estimating confidence limits (14). Rare species, with $N < 50$ individuals, had to be dealt with carefully, and most of our statistics are based only on species with at least one individual per hectare (15).

Nearly every species was aggregated when all diameter classes of ≥ 1 cm were included. In the six plots, 1768 species had at least one individual per hectare: 1753 were aggregated at 0 to 10 m ($\Omega_{0-10} > 1$), 1490 significantly so (95% confidence limits around Ω_{0-10} did not include 1); 1759 were aggregated at 10 to 20 m, and 1730 were aggregated at 20 to 30 m (16). No plot had fewer than 96% of its species aggregated at 0 to 10 m, and every one of 772 species at Lambir was aggregated. Relative neighborhood density almost invariably declined with distance (Fig. 1): $\Omega_{10-20} < \Omega_{0-10}$ in 1714 of 1768 species, and $\Omega_{20-30} < \Omega_{10-20}$ in 1581 species. Nevertheless, Ω values in nearby distance classes were highly correlated with one another. Thus, we can use Ω_{0-10} —the mean conspecific density within 10 m of a tree (relative to the species' overall density)—as a simple measure of the intensity of aggregation of a species.

There was an enormous range in Ω_{0-10} . Most species had values of <10 , but there were a few species with much higher aggregation. The highest of all among the species with ≥ 1 individual per hectare, $\Omega_{0-10} = 906$, was in *Lagerstroemia* sp. (Lythraceae) in the HKK plot; 51 of the 59 individuals of this species occurred in a single clump in an area of 400 m². Nineteen species had aggregation indices over 100, eight at Lambir and four at HKK. These results are dominated by saplings, because the vast majority of

trees ≥ 1 cm diameter are <10 cm diameter.

Results hold for larger trees, though. Nearly all species were aggregated when only diameter classes of ≥ 10 cm were considered. There were 543 species with more than 1 individual per hectare at this size in all six plots, and 488 had $\Omega_{0-10} > 1$; for 257 species, the aggregation was significant (16). But aggregation intensity weakened at greater sizes in most species: 321 of 543 had a higher Ω_{0-10} for all trees than for larger trees and of those with a significant difference, 84 of 102 were more aggregated at the smaller than at the larger size. At BCI, Pasoh, HKK, and Lambir, about two-thirds of the species were more aggregated at the smaller diameter class, but the pattern reversed at Sinharaja and Mudumalai and most species became more aggregated at the larger size. We repeated the analysis for trees of ≥ 30 cm diameter and again found most species aggregated (16).

Rare species were substantially more aggre-

gated than common species at all but the Mudumalai site (17). Median Ω_{0-10} was 4 to 10 times higher in the rare abundance class than in the commonest classes (Table 2). In the most abundant species, $\Omega_{0-10} < 5$, and at BCI and Pasoh mostly $\Omega_{0-10} < 2$, whereas in species with fewer than 500 individuals, Ω_{0-10} typically ranged from 5 to 30 or higher (Fig. 2). This trend held for species as rare as 10 individuals per 50 ha, even though many species had $\Omega_{0-10} = 0$ when $N \cong 10$ (15). For $N < 10$ individuals, median scores were zero, but arithmetic means were very high. Thus, the rarest species were the most aggregated of all, and across the entire range of abundances (four orders of magnitude), the degree of clumping correlated negatively with species density. These results match Hubbell's conclusion from dry forest in Costa Rica (4) and two previous analyses of the Pasoh plot (18, 19). All results from the neighborhood density statistic Ω were

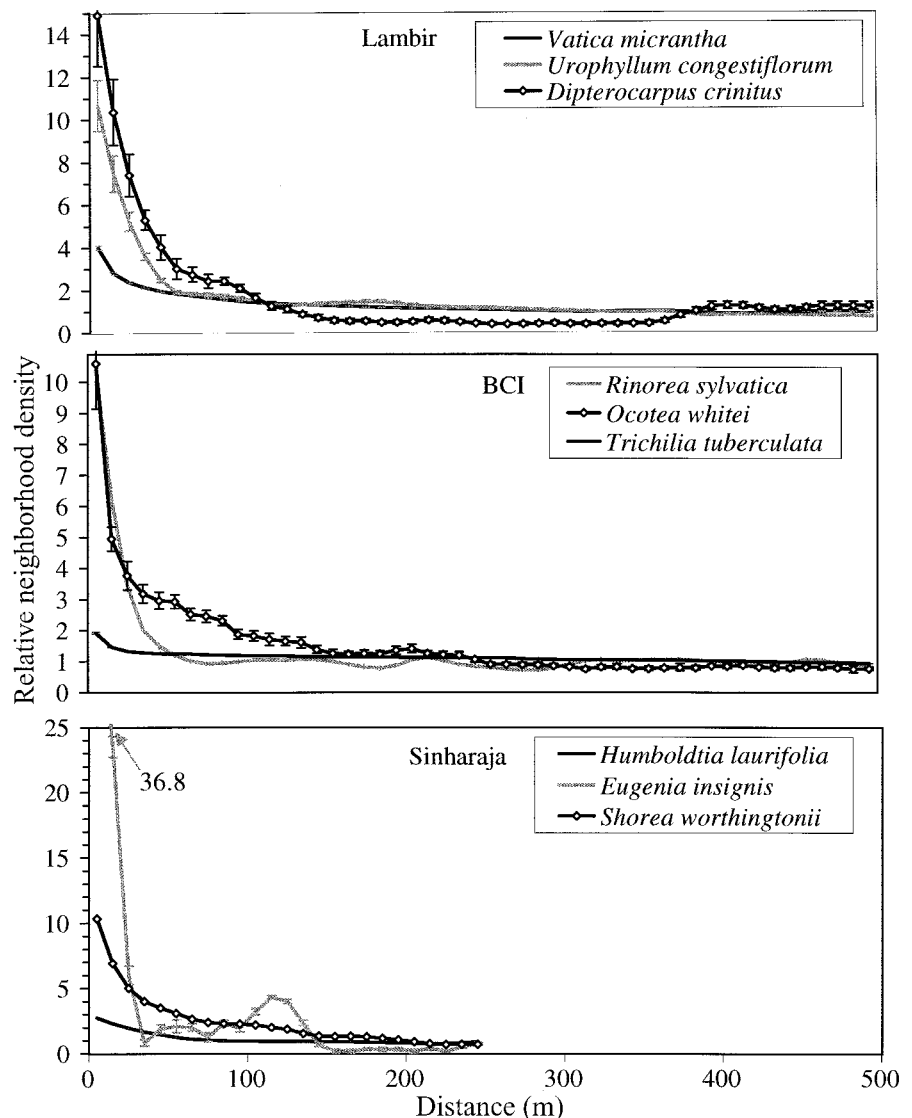


Fig. 1. Relative neighborhood density (Ω_x as defined in the text) as a function of distance x in sample species. Vertical bars give 95% confidence limits (14), sometimes too small to see.

R E P O R T S

Table 1. Six tropical forest dynamics plots that have been fully censused at least once. Underlined census year is the one used in this report. Dry season months gives the number per year with mean rainfall < 100 mm. dbh, diameter at breast height.

Plot	Plot size (ha)	Fully censused	Topographic variation (m)	Annual rainfall (mm)	Dry season (months)	Species ≥ 1 cm dbh	Species ≥ 10 cm dbh	Individuals ≥ 1 cm dbh	Individuals ≥ 10 cm dbh
Barro Colorado Island (BCI), Panama	50	1981–83 1985 1990 1995	38	2500	4	299	226	229,071	21,459
Pasoh, Malaysia (Peninsula)	50	1986–89 1990 1995–96	25	1800	0	818	673	320,382	28,997
Lambir, Malaysia (Borneo)	52	1991–94 1997	140	2700	0	1174	996	366,121	32,962
Huai Kha Khaeng (HKK), Thailand	50	1992–94	89	1450	6	248	211	81,145	21,957
Mudumalai, India	50	1988–89 1992 1996	130	1200	6	72	63	25,306	14,922
Sinharaja, Sri Lanka	25	1996–98	151	5000	0	203	164	206,227	17,214

Table 2. Median aggregation index, Ω_{0-10} , across species in various abundance categories in the six large plots. Number of species within each abundance category is listed under spp. Last row gives overall median Ω_{0-10} for all species with at least 50 individuals at each plot.

Abundance class (per 50 ha)	Pasoh		BCI		Mudumalai		Sinharaja		Lambir		HKK	
	Median Ω_{0-10}	spp.	Median Ω_{0-10}	spp.	Median Ω_{0-10}	spp.	Median Ω_{0-10}	spp.	Median Ω_{0-10}	spp.	Median Ω_{0-10}	spp.
≥ 5000	1.5	7	1.5	7	4.4	1	3.7	19	4.0	3	1.5	3
2000–4999	1.8	22	2.2	16	2.7	5	3.3	20	3.6	22	2.5	7
1500–1999	1.8	20	1.7	10	2.7	1	3.6	11	2.7	12	2.9	6
1000–1499	2.1	30	2.3	12		0	4.2	12	2.5	34	3.1	5
750–999	2.0	34	2.5	11	10.3	1	4.1	6	3.7	37	3.0	7
500–749	2.3	58	2.6	19	1.6	2	3.7	16	3.7	72	5.9	8
300–499	2.7	76	2.7	18	2.7	2	4.7	28	4.1	121	2.4	6
200–299	3.1	73	4.0	20	3.1	1	7.3	14	5.3	111	11.7	9
150–199	3.2	51	3.3	11	3.1	1	13.9	6	5.4	78	5.1	14
100–149	4.0	68	4.0	23	29.6	3	9.5	9	7.0	118	11.9	10
75–99	5.4	48	6.3	18	11.4	5	13.7	9	9.1	74	5.4	8
50–74	6.1	48	10.9	18	1.7	3	12.6	8	11.5	104	6.1	12
25–49	8.3	81	7.8	22	8.9	11	15.8	16	14.0	148	13.3	26
10–24	14.5	89	22.1	27	27.7	12	0.0	8	14.3	112	3.0	32
≥ 50	3.0	535	3.1	183	3.4	25	4.7	161	5.6	772	4.6	95

confirmed by nearest-neighbor analysis (16).

The six forests were remarkably similar in community-wide patterns of aggregation. The range of values for Ω_{0-10} was similar at all sites, and the relationship between this aggregation index and abundance was essentially identical across sites. However, there were significant differences (20). Pasoh and BCI had low aggregation indices at a given abundance, whereas Sinharaja and Lambir had high indices (Table 2). These results correspond with habitat variation within the plots in that Pasoh and BCI are topographically uniform and Lambir and Sinharaja are relatively rugged (Table 1). Lambir also has a sharp soil gradient (21). Mudumalai has steep topography, but there is little indication that species respond to it, whereas at Lambir and Sinharaja many species' distributions follow topographic features (22). There are clear exam-

ples of habitat-related patchiness at most of the plots, especially Sinharaja (Fig. 3). Species in which larger trees were more densely aggregated than juveniles are also suggestive of habitat-related patchiness, because adults might be collected in sites most favorable for the species, whereas juveniles are widely dispersed.

On the other hand, there are many species in all the plots whose aggregated distributions indicate dispersal limitation. These species occur in circular clumps that do not correspond with topography (Fig. 3), and they had the highest values of Ω_{0-10} . Their neighborhood density functions Ω_x declined abruptly with distance (Fig. 1). We tested the importance of dispersal limitation by comparing aggregation intensity in poorly versus well-dispersed trees. Species whose seeds are dispersed by animals were assumed to be better

dispersed than wind- or explosively dispersed species, and canopy trees were assumed to have well-dispersed seeds relative to understory treelets (23). We also considered the Dipterocarpaceae, a family of trees with poorly dispersed, winged seeds that dominates Southeast Asian forests.

The prediction that better dispersal reduces aggregation was partially borne out. There was no significant difference in aggregation intensity between canopy and understory species at Pasoh, but at BCI there was (24). At BCI, there was no significant difference in aggregation for animal versus nonanimal dispersed species, but the difference was fairly pronounced in the predicted direction (25). Finally, at the two Malaysian plots, dipterocarps were strikingly more aggregated than nondipterocarps (26).

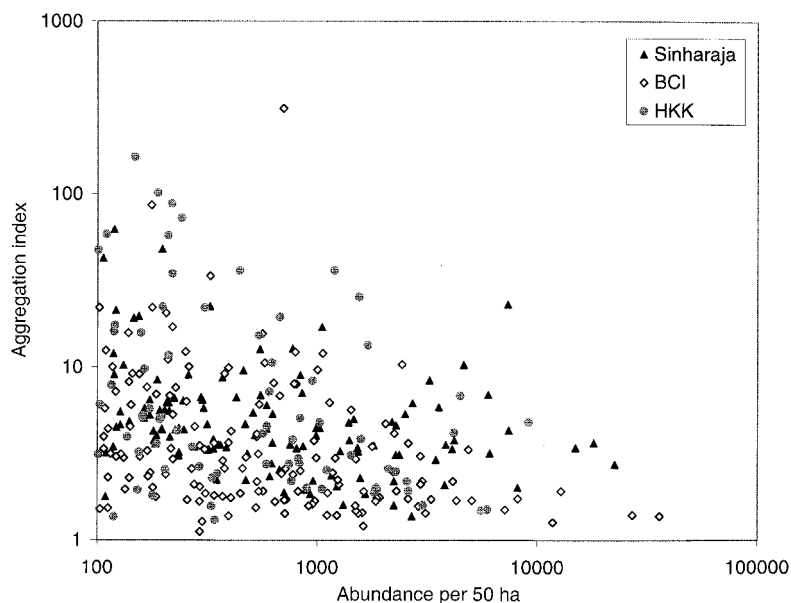


Fig. 2. Aggregation index (Ω_{0-10} , the relative density of conspecifics within 10 m of focal trees) for all species with ≥ 100 individuals at three plots, as a function of the abundance of each species, on a log-log scale.

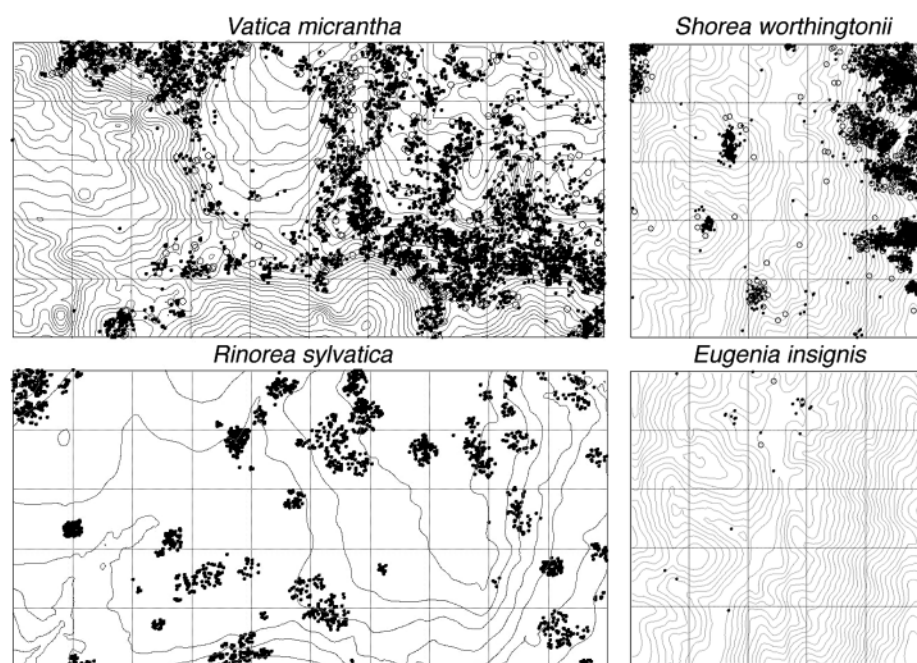


Fig. 3. Distribution maps for species also used in Fig. 1. Small circles, trees of 1 to 9.9 cm diameter; open circles, trees of ≥ 10 cm diameter. Grid squares = 1 ha. *Vatica* clumps follow ridges at Lambir. *Rinorea* clumps at BCI do not correlate with any known canopy, topographic, or soil feature, and the patches are probably due to limited seed dispersal (seeds disperse from exploding capsules). *Shorea* follows ridge tops at Sinharaja, and *Eugenia* is very rare at Sinharaja, but most individuals are close to several conspecifics. Additional maps published elsewhere (32, 33) illustrate many cases of habitat and dispersal limited patchiness.

Finally, the observation that aggregation is weaker in larger diameter classes supports the notion that herbivores and plant diseases play a role in reducing aggregation. Other evidence from the BCI and Pasoh plots indicates that pests have already substantially

weakened aggregation intensity by the time trees enter the census at 1 cm diameter (27, 28). After 1 cm diameter, the pest effect is not particularly dramatic, and we saw no indication for further loosening of aggregations between 10- and 30-cm diameters.

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15. Even under complete spatial randomness, the probability that $\Omega_{0-10} = 0$ is high when $N < 50$ in 50 ha. Across all plots, no species with $N \geq 100$ had $\Omega_{0-10} = 0$, whereas a few species with 50 to 100 individuals had $\Omega_{0-10} = 0$. With $N < 10$, $\Omega_{0-10} = 0$ in most species. For this reason, most of our analyses refer to all species with ≥ 50 individuals in the 50-ha plots (to standardize density, the cutoff was set at 1 per hectare, or 25 individuals in the Sinharaja plot and 52 in the Lambir plot).
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24. At Pasoh, the median Ω_{0-10} for canopy species was

- 2.8, and for understory it was 3.1; the difference is not significant. At BCI, the medians were 2.9 and 4.2, and the difference is significant.
25. The median Ω_{0-10} for wind- or explosively dispersed species at BCI was 4.5; for animal-dispersed species it was 3.0, but the difference was not significant.
26. At Pasoh, the 24 dipterocarp species with $N \geq 50$ individuals had a median Ω_{0-10} of 5.1, compared with 2.8 for all other species. At Lambir, 65 dipterocarps had $\Omega_{0-10} = 21.8$; for other species, $\Omega_{0-10} = 5.3$. Both differences are significant. At Sinharaja, 12 dipterocarps were slightly but not significantly more aggregated than nondipterocarps. HKK and Mudumalai had just three and one dipterocarp species, respectively.
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Mechanism of ATP-Dependent Promoter Melting by Transcription Factor IIH

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We show that transcription factor IIH ERCC3 subunit, the DNA helicase responsible for adenosine triphosphate (ATP)-dependent promoter melting during transcription initiation, does not interact with the promoter region that undergoes melting but instead interacts with DNA downstream of this region. We show further that promoter melting does not change protein-DNA interactions upstream of the region that undergoes melting but does change interactions within and downstream of this region. Our results rule out the proposal that IIH functions in promoter melting through a conventional DNA-helicase mechanism. We propose that IIH functions as a molecular wrench: rotating downstream DNA relative to fixed upstream protein-DNA interactions, thereby generating torque on, and melting, the intervening DNA.

Human transcription factor IIH consists of nine polypeptides with masses of 31 to 90 kD (1-3). IIH is responsible for three critical functions in transcription: phosphorylation of the COOH-terminal domain (CTD) of the RPB1 subunit of RNA polymerase II (RNAPII), promoter melting, and promoter clearance.

IIH-dependent CTD phosphorylation re-

quires ATP and is mediated by the IIH cdk7 subunit, which is a cyclin-dependent protein kinase (1-3). The role of IIH in promoter melting is to melt about one turn of DNA encompassing the transcription start site to yield the "transcription bubble" (4, 5). This process requires ATP (4, 5) and is mediated by the IIH ERCC3 subunit (6, 7) (also referred to as

XPB), which, in isolation, exhibits 3'-5' DNA-helicase activity (6, 8, 9). The role of IIH in promoter clearance is to stimulate escape of transcription elongation complexes stalled at positions +10 to +17 (10-12). Like promoter melting, promoter escape requires ATP (10-13) and is mediated by the IIH ERCC3 subunit (14).

The fact that promoter melting involves generation of single-stranded DNA (ssDNA) and that promoter escape involves a species containing ssDNA, together with the fact that the IIH subunit that mediates these processes exhibits DNA-helicase activity, has led to the proposal that IIH functions in these processes through a conventional DNA-helicase mechanism, with direct interactions between IIH and ssDNA (1-3, 15). However, no direct evidence has been obtained in support of this proposal.

As a first step to understand the mechanism

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Fig. 1. Results of protein-DNA photo-cross-linking experiments. **(A)** Representative data. TBFR, transcription-complex intermediate containing RNAPII, TBP, IIB, IIF, and promoter DNA (78); TBFRE, TBFR plus IIE; TCC, transcriptionally competent complex, consisting of TBFR plus IIE and IIH; TCC + ATP, transcriptionally competent complex after ATP-dependent CTD phosphorylation and promoter melting. RPB1 and RPB1-Pn denote forms of the largest subunit of RNAPII having unphosphorylated CTD and phosphorylated CTD, respectively. Data are shown for positions -2, +5, +13, and +21 of the DNA nontemplate strand. **(B)** Representative data for experiments assessing NTP specificity (positions -2, +5, and +13 of DNA nontemplate strand). **(C)** Representative data confirming identities of cross-linked IIH α [(left) parallel experiment with IIH α (1-394) (31, 49)] and IIH ERCC3 [(right) immunoprecipitation with antibody to ERCC3 of cross-linked polypeptide (32)]. **(D)** Representative data demonstrating increase in fraction of complexes competent for CTD phosphorylation (top; analysis of electrophoretic mobility of cross-linked RPB1) and transcription [bottom; quantitation of transcription (arbitrary units) and RNAPII content (1.0 unit = 20 fmol RNAPII) upon Sarkosyl washing (24)].

