Background: The dynamics of subunit incorporation in complex I (CI) under steady-state conditions are still elusive.

Results: GFP-labeled CI subunits replace their endogenous counterparts in CI with different efficiencies and rates within 24 h of induction.

Conclusion: CI maintenance involves continuous subunit replacement by direct exchange or through assembly intermediates.

Significance: Elucidating subunit replacement in CI is vital as subunits are continuously threatened by oxidative damage.

Studies employing native PAGE suggest that most nDNA-encoded CI subunits form subassemblies before assembling into holo-CI. In addition, in vitro evidence suggests that some subunits can directly exchange in holo-CI. Presently, data on the kinetics of these two incorporation modes for individual CI subunits during CI maintenance are sparse. Here, we used inducible HEK293 cell lines stably expressing AcGFP1-tagged CI subunits and quantified the amount of tagged subunit in mitoplasts and holo-CI by non-native and native PAGE, respectively, to determine their CI incorporation efficiency. Analysis of time courses of induction revealed three subunit-specific patterns. A first pattern, represented by NDUFS1, showed overlapping time courses, indicating that imported subunits predominantly incorporate into holo-CI. A second pattern, represented by NDUFV1, consisted of parallel time courses, which were, however, not quantitatively overlapping, suggesting that imported subunits incorporate at similar rates into holo-CI and CI assembly intermediates. The third pattern, represented by NDUFS3 and NDUFA2, revealed a delayed incorporation into holo-CI, suggesting their prior appearance in CI assembly intermediates and/or as free monomers. Our analysis showed the same maximum incorporation into holo-CI for NDUFV1, NDUFV2, NDUFS1, NDUFS3, NDUFS4, NDUFA2, and NDUFA12 with nearly complete loss of endogenous subunit at 24 h of induction, indicative of an equimolar stoichiometry and unexpectedly rapid turnover. In conclusion, the results presented demonstrate that newly formed nDNA-encoded CI subunits rapidly incorporate into holo-CI in a subunit-specific manner.

Mitochondrial NADH:ubiquinone oxidoreductase or complex I (CI; EC 1.6.5.3) is the largest enzyme of the oxidative phosphorylation system. It catalyzes the oxidation of NADH and reduction of ubiquinone, utilizing the energy generated by this process to translocate protons across the mitochondrial inner membrane (1). Human CI consists of at least 44 different subunits (2), with seven encoded by the mitochondrial DNA (mtDNA) and the remainder by the nDNA. The manner in which this intricate enzyme is maintained is only beginning to be understood.

According to the current view on CI assembly, subunits of the dehydrogenase, hydrogenase, and proton-translocation module assemble via several intermediates into the L-shaped holoenzyme (3–9). In addition to the de novo CI assembly pathway, a CI subunit exchange pathway was proposed, in which specific subunits (NDUFV1, NDUFV2, NDUFV3, NDUFS6, and NDUFS4) directly replace their preexisting counterparts in holo-CI, whereas other subunits (NDUF8, NDUFS1, NDUFS2, NDUFS7, NDUFS8, NDUFA9, NDUFA10) lack this capability (8) (see Table 1 for subunit nomenclature in different species). However, the extent to which these two incorporation manners are applied under physiological conditions remains to be elucidated.

Although a basic framework for human CI assembly has been established, the turnover and assembly dynamics of individual CI subunits under non-disturbed, steady-state conditions remain largely unknown. Pulse labeling studies in the fungus Neurospora crassa revealed that newly synthesized subunits assemble at different rates into holo-CI (10). This suggests that not all CI subunits are imported, assembled, exchanged, and/or maintained at similar rates. Only few studies have investigated the assembly dynamics of mammalian CI (3, 8, 11–14). Pulse labeling studies established that, in general, nDNA-encoded subunits appear faster in holo-CI than mtDNA-encoded subunits (8, 11). After complete depletion of CI by transient inhibition of mitochondrial translation, osteosarcoma cells needed 2–4 days to normalize their CI level (3, 12, 14). A similar restoration time was observed after inducible HA-tagged NDUFA1 expression in NDUFA1-deficient Chinese hamster cells (13). In molecular weight; NDUF, NADH dehydrogenase ubiquinone flavoprotein; IGA, in-gel activity; EGFP, enhanced GFP.

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these mutant Chinese hamster fibroblasts, holo-CI formation clearly lagged behind the increase in NDUFA1-HA, suggesting that this subunit first appears in a precomplex, without using an exchange route in parallel (13). However, the dynamics of subunit incorporation during these CI depletion/restoration experiments are very likely to differ from the natural assembly kinetics during undisturbed CI maintenance because (i) direct subunit exchange in non-existing CI is per definition not possible and (ii) the pool sizes of individual subunits and subassemblies alter upon CI depletion (3, 4, 13, 15–19).

There are still many aspects of mammalian CI homeostasis that remain to be elucidated. For instance, no definite data are available on (i) the incorporation efficiency of imported CI subunits into holo-CI, (ii) the stoichiometry of CI subunits in holo-CI, and (iii) the incorporation kinetics of individual CI subunits.

In this study, we addressed these topics using inducible human CI, and (i) the incorporation kinetics during undisturbed CI maintenance because (i) direct subunit exchange in non-existing CI is per definition not possible and (ii) the pool sizes of individual subunits and subassemblies alter upon CI depletion (3, 4, 13, 15–19).

**TABLE 1**

Nomenclature of CI subunits investigated in this study

| Hs | Br | Yl | Tt | Ec |
|----|----|----|----|----|
| NDUFV1 | 51-kDa | NUBM | Nqo1 | NuoF |
| NDUFV2 | 24-kDa | NUTH | Nqo2 | NuoE |
| NDUFS1 | 75-kDa | NUAM | Nqo3 | NuoG |
| NDUFS2 | 49-kDa | NUCM | Nqo4 | NuoCD |
| NDUFS3 | 30-kDa | NUGM | Nqo5 | NuoCD |
| NDUFS7 | PSSV | NUKM | Nqo6 | NuoB |
| NDUFS8 | TYKY | NUIM | Nqo9 | NuoL |
| NDUFS4 | AQDQ | NUYM | n.p. | n.p. |
| NDUFS6 | 13-kDa | NUNM | n.p. | n.p. |
| NDUFA2 | 48 | NIBM | n.p. | n.p. |
| NDUFA12 | B17.2 | N7BM | n.p. | n.p. |
| NDUFA1 | MWFE | NIMM | n.p. | n.p. |
| NDUFB6 | B17 | NPM | n.p. | n.p. |
| NDUFB8 | ASHI | NIAM | n.p. | n.p. |

**EXPERIMENTAL PROCEDURES**

*Generation and Culturing of Inducible Cell Lines—Gateway™*

Entry vectors (Invitrogen) containing CI subunit open reading frames without stop codon (NDUFV1, NM_007103; NDUFS1, NM_005006; NDUFS2, NM_004550; NDUFS3, NM_004551; NDUFS7, BC 111517; NDUFS8, NM_002496; NDUFS4, NM_002495; NDUFS6, BC_046155; NDUFA2, NM_002488; NDUFA12, NM_018838; NDUFA1, NM_004541; NDUFB6, NM_002493; transcript variant 1) or the COX8 mitochondrial targeting sequence (first 210 base pairs of sequence NM_00004074) were created as described previously (19). Entry clones containing NDUFV2 (BC_001632; clone HsCD0000415552), NDUFB8 (BC_000466; clone HsCD000042304) without stop codon were ordered from The Plasmid Database (22). A destination vector containing the AcGFP1 sequence (Clontech, Westburg, Leusden, The Netherlands) in-frame behind Gateway Reading Frame Cassette B (Invitrogen) and a Flp Recombination Target site for targeted genomic integration in Flp-in T-Rex293 cells was created as described previously (19). Expression clones were generated by recombining the entry clones with the destination vector by using Gateway LR Clonase II enzyme mix (Invitrogen) (for details, see Ref. 19). Expression vectors were co-transfected with the pOG44 vector (Invitrogen) in Flp-in T-Rex293 cells (Invitrogen) using Superfect transfection Reagent (Qiagen, Venlo, The Netherlands). Cells were cultured for selection in DMEM (BioWhitaker, Walkersville, MD) supplemented with 10% (v/v) fetal calf serum (FCS) (PAA Laboratories, Pasching, Austria), 1% penicillin/streptomycin (Invitrogen), 200 μg/ml hygromycine (Calbiochem, Brunschwig, Amsterdam, The Netherlands), and 50 μg/ml blasticin (Invitrogen). For induction of overexpression, doxycycline was added at a concentration of 1 μg/ml for 24 hr.

**Quantitative Real-time PCR—RNA was isolated from cell pellets from two independent cultures using TRizol reagent (Invitrogen) according to the manufacturer’s instructions. RNA was dissolved in RNase-free water at 65 °C and diluted to a final concentration of 200 ng/μl. Reverse transcription was performed in a total mixture of 40 μl containing 2 μg of RNA, 1× first strand buffer (Invitrogen), 0.01 μM DTT (Invitrogen), 1 mM dNTP mix, 200 units of Superscript II reverse transcriptase (Invitrogen), 60 units of RNAsin (Promega, Madison, WI), 625 ng oligo(dT) primers (Promega), and 125 pmol of random primers (Promega). This reaction mixture was incubated at 42 °C for 1 h and at 96 °C for 3 min. The cDNA samples of two independent experiments were analyzed on two different days by quantitative real-time PCR. Separate real-time PCR reactions were performed for AcGFP1, β-actin, and 18S ribosomal RNA (18 S rRNA) on 50 ng of cDNA using SybGreen PCR Master Mix (Applied Biosystems, Zwijndrecht, The Netherlands), 15 pmol of forward primer and 15 pmol of reverse primer in a reaction volume of 25 μl. Primers were designed by PrimerExpress Software (Applied Biosystems). The primer pairs used were as follows: AcGFP1_fwd 5′-gaggatgacggcaactaatggc-3′ with AcGFP1_rev, 5′-acacattgcccgtttgattg-3′; β-actin_fwd, 5′-cgattgcaacctgcagacagac-3′ with β-actin_rev, 5′-ctaggagaccagatgtctgg-3′; 18 S rRNA_fwd, 5′-GTAACCGTGTGAACCCATT-3′ with 18 S rRNA_rev, 5′-CCATCCCAATCGGATGTAGCCG-3′. Thermal cycling conditions were set as follows: 2 min at 50 °C and 10 min at 95 °C, followed by 40 cycles of 15 s at 95 °C and 1 min at 60 °C. Amplification of a single product for each primer set was first confirmed by electrophoresis on a 2% agarose gel. PCR amplification and detection was performed using an ABI Prism 7900 HT Sequence Detection System (Applied Biosystems). A comparative threshold cycle (Ct) method was used to quantify the gene expression of AcGFP1-tagged CI subunits in different cell lines. As a calibrator, we analyzed AcGFP1-expression in a cell line expressing matrix-targeted AcGFP1 (named GFP). To compare the expression in different cell lines, the Ct of β-actin was subtracted from the Ct of GFP to yield the ΔCt. The expression of AcGFP1-tagged CI subunits was expressed as 2−ΔΔCt, where ΔΔCt = (ΔCt_GFP – ΔCt_AcGFP1-tagged CI subunit). Normalization to 18 S rRNA was performed in a similar way to confirm results obtained using β-actin. Statistical analysis was performed using Origin Pro (version 7.5, Originlabs, Northampton, MA). Values are expressed as mean ± S.D.
BN-PAGE and SDS-PAGE Analysis—Mitoplasts were isolated from cell pellets using digitonin, and mitochondrial protein complexes were solubilized using n-dodecyl β- d-maltoside as described previously (23). BN-PAGE 5–15% gradient gels were loaded with 40–80 μg protein and scanned for fluorescence (473 excitation laser, FITC filter/510 nm) using a FLA5000 fluorescence image analyzer (Fujifilm, Tilburg, The Netherlands). Subsequently, CI in-gel activity (IGA) assays were performed, or proteins were blotted to a nitrocellulose membrane. A small fraction of the mitochondrial lysates was denatured and loaded on 10% SDS-PAGE gels as described previously (4), scanned for fluorescence and blotted to nitrocellulose. Immunodetection was performed using primary antibodies raised against EGFP (a gift from Dr. F. J. van Kuppeveld, Nijmegen, the Netherlands), succinate dehydrogenase complex subunit A (Invitrogen), NDUFV2 (Sigma, Zwijndrecht, The Netherlands), NDUF52 (a gift from Prof. B. Robinson, Toronto, Canada), NDUF53 (Invitrogen), NDUF54 (Mitosciences, Eugene, OR), NDUFA2 (4), NDUFA1 (a gift from Prof. I. Scheffler, San Diego, CA), and NDUFB6 (Invitrogen). Secondary antibodies were peroxidase-conjugated anti-mouse or anti-rabbit IgGs (Invitrogen). Chemiluminescence (Pierce, Thermo Fisher Scientific, Etten-Leur, The Netherlands) was detected on film or on a LAS-3000 luminescent image analyzer (Fujifilm).

RESULTS

Incorporation of AcGFP1-tagged CI Subunits into Holo-CI—To study the incorporation kinetics of nDNA-encoded subunits into holo-CI, we used a panel of Flp-In T-Rex293 cell lines engineered to express AcGFP1-tagged versions of these subunits in a doxycycline-inducible manner (4, 19, 20). In the remainder of the paper, the prefix “t” is used to indicate the tagged version of the subunit in question. Apart from the seven nDNA-encoded core subunits (NDUFV1, NDUFV2, NDUFS1, NDUFS2, NDUFS3, NDUFS7, and NDUFS8) we also investigated seven accessory subunits (NDUF54, NDUFS6, NDUFA1, NDUFA2, NDUFA12, NDUFB6, and NDUFB8), predicted to constitute part of either the peripheral or membrane arm of CI (Fig. 1A).
**Dynamics of Mitochondrial Complex I Assembly**

In theory, the cellular transcript levels of the tagged CI subunits should be the same because we used targeted recombination, meaning that cDNAs were integrated at the same position in the genomic DNA and lacked the 5’ and 3’ post-transcriptional regulatory elements (Fig. 1B). Indeed, quantitative RT-PCR analysis revealed no significant difference in the amount of transcripts between the various cell lines (Fig. 1C). BN-PAGE of mitoplast fractions followed by Western blotting with an anti-GFP antibody showed that all cell lines expressed the AcGFP1-tagged subunit (Fig. 1D). Optimal exposure of each individual lane revealed that the tagged CI subunits were generally incorporated in holo-CI (filled arrowhead) and in some cases also in lower MW subcomplexes. Exceptions were tNDUFS2 and tNDUFS7, which were clearly detectable in lower MW subcomplexes but hardly in holo-CI.

Mitoplast Accumulation and Holo-CI Incorporation of AcGFP1-tagged CI Subunits at a High Level of Induction—Expression of tagged subunits was increased by culturing the cells in the presence of 1 μg/ml doxycycline for 24 h. Next, we isolated mitoplasts, separated their constituent proteins by SDS-PAGE, and determined the total amount of tagged subunit. Fluorescence scanning of the obtained gels demonstrated the presence of both intact tagged subunit (filled arrowheads) and breakdown products with a lower MW (Fig. 2A). To assess the amount of tagged subunit incorporated into holo-CI, we analyzed the same mitoplast preparations using BN-PAGE. Comparison of the resulting BN-PAGE fluorogram (Fig. 2B) with the immunoblot depicted in Fig. 1D revealed a similar pattern. Next, the BN-PAGE gel was used for IGA analysis, revealing that all cell lines displayed an IGA-positive band at the height of holo-CI (Fig. 2C). The intensity of this band was similar among the various cell lines and not affected by expression induction. This is consistent with our previous studies demonstrating that overexpression of tNDUFV1, tNDUFV2, tNDUFS3, tNDUFA2, tNDUFA12, and tNDUF6 does not interfere with CI activity. Comparison of the resulting BN-PAGE fluorogram (Fig. 2B) with the immunoblot depicted in Fig. 1D revealed a similar pattern. Next, the BN-PAGE gel was used for IGA analysis, revealing that all cell lines displayed an IGA-positive band at the height of holo-CI (Fig. 2C). The intensity of this band was similar among the various cell lines and not affected by expression induction. This is consistent with our previous studies demonstrating that overexpression of tNDUFV1, tNDUFV2, tNDUFS3, tNDUFA2, tNDUFA12, and tNDUF6 does not interfere with CI activity. Comparison of the resulting BN-PAGE fluorogram (Fig. 2B) with the immunoblot depicted in Fig. 1D revealed a similar pattern. Next, the BN-PAGE gel was used for IGA analysis, revealing that all cell lines displayed an IGA-positive band at the height of holo-CI (Fig. 2C). The intensity of this band was similar among the various cell lines and not affected by expression induction. This is consistent with our previous studies demonstrating that overexpression of tNDUFV1, tNDUFV2, tNDUFS3, tNDUFA2, tNDUFA12, and tNDUF6 does not interfere with CI activity. Comparison of the resulting BN-PAGE fluorogram (Fig. 2B) with the immunoblot depicted in Fig. 1D revealed a similar pattern. Next, the BN-PAGE gel was used for IGA analysis, revealing that all cell lines displayed an IGA-positive band at the height of holo-CI (Fig. 2C). The intensity of this band was similar among the various cell lines and not affected by expression induction. This is consistent with our previous studies demonstrating that overexpression of tNDUFV1, tNDUFV2, tNDUFS3, tNDUFA2, tNDUFA12, and tNDUF6 does not interfere with CI activity.

**Incorporation Efficiency of AcGFP1-tagged CI Subunits into Holo-CI at a Low Level of Induction**—Next, we analyzed the amount of tagged subunit present in mitoplasts (Fig. 5A) and holo-CI (Fig. 5B) of non-induced cells. These cells displayed a low steady-state (“leakage”) expression of the tagged subunit due to small amounts of tetracycline in the culture medium. Using an identical analysis as for Fig. 2D, it was found that a maximum incorporation efficiency was reached for tNDUFA1, tNDUF4, tNDUFS1, and tNDUFV2 (Fig. 5D, data points on the line of identity). In contrast, low efficiencies were reached for tNDUFA12 and tNDUF6 (data points well below the line of identity).

**Temporal Dynamics of Mitochondrial Import and CI Incorporation of AcGFP1-tagged CI Subunits**—To determine the dynamics of subunit increase in mitoplasts and holo-CI, we investigated the amount of tagged subunit during the transition from low- to high-level expression. For this analysis, we used four key subunits (NDUFV1, NDUF51, NDUF53, and NDUFA2) that reached maximum incorporation into holo-CI within 24 h after induction and showed similar incorporation...
FIGURE 2. Incorporation efficiency of AcGFP1-tagged Cl subunits after induction with doxycycline. A, cells were cultured in the presence of 1 μg/ml doxycycline for 24 h to obtain high steady-state expression. Identical protein amounts of isolated mitoplasts were subjected to SDS-PAGE, and gels were scanned for fluorescence. Bands of predicted MW are indicated with filled arrowheads. The first lane is from cells expressing matrix-targeted AcGFP1 (marked GFP) after addition of two MW marker proteins (marked M25 and M75) to the mitoplast sample. Lower MW bands reflect breakdown products (see also Ref. 20). B, identical protein amounts of the above mitoplast preparations were separated by BN-PAGE and scanned for fluorescence. The position of holo-CI is indicated with a filled arrowhead. C, the same gel as in B after staining for in-gel CI activity (IGA). Of note, compared with the gel in B, the right part of the gel in C is bent down because of the procedure for IGA measurement. D, incorporation efficiency plot of CI subunits. The fluorescence intensity of intact AcGFP1-tagged subunit on the SDS-PAGE fluorogram (A) was normalized to the corresponding IGA signal (C) and expressed as a percentage of the value obtained with intact matrix-targeted AcGFP1 (x axis). Similarly, the fluorescence intensity of holo-CI on the BN-PAGE fluorogram (B) was normalized to the corresponding IGA signal (C) and expressed as a percentage of the value obtained with intact matrix-targeted AcGFP1 (y axis). Thus, the x axis reflects the total amount of intact tagged subunit present in the mitoplasts, whereas the y axis reflects the amount of tagged subunit incorporated in holo-CI. The solid line (line of identity) indicates maximum incorporation efficiency, whereas the dashed line represents the maximum incorporation level.
efficiencies at low and high levels of steady-state expression (Fig. 6). Cells were harvested at various time points after induction and mitoplast proteins were separated by SDS- and BN-PAGE (for a typical example of such an experiment, see Fig. 1 of our previous study (4)). At each time point, we quantified the total amount of tagged subunit in the mitoplast or in holo-CI and normalized it to the corresponding IGA band. Values determined at 24 h were either set to the values in Fig. 2D (Fig. 6) or to 100% (Fig. 6; insets). The first normalization allows evaluation of the levels of tagged subunit in mitoplasts and holo-CI, whereas the second normalization enables better comparison of their import and incorporation kinetics.

For all tagged subunits, except tNDUFV1, mitoplast accumulation (solid line) and holo-CI incorporation (dotted line) reached a plateau within 24 h of induction. The time courses for mitoplast entry and holo-CI incorporation largely coincided for tNDUS1 and tNDUFV1. This indicates that, once imported, incorporation of these subunits occurs rapidly. In contrast, for tNDUS3 and tNDUF2 CI incorporation clearly lagged behind their accumulation in the mitoplasts.

DISCUSSION

Data from previous studies with cells with a naturally or pharmacologically disturbed CI assembly have been interpreted to indicate that CI de novo synthesis occurs via several steps, in which subunits from different functional modules first form smaller assembly intermediates that are then combined to constitute the holoenzyme (3, 4, 6–9, 20). In addition, in vitro
Incorporation efficiency of AcGFP1-tagged CI subunits into holo-CI at low steady-state expression. Cells were cultured in normal growth medium containing traces of tetracycline from FCS causing leakage expression. A, from each cell line, identical amounts of isolated mitoplasts were subjected to SDS-PAGE and scanned for fluorescence. Bands of predicted MW are indicated with filled arrowheads. The first lane is from mitoplasts of cells expressing matrix-targeted AcGFP1 (marked GFP) after addition of two MW marker proteins (marked M25 and M75). B, identical amounts of the same mitoplast preparations were separated by BN-PAGE and scanned for fluorescence. Filled arrowheads indicate the position of holo-CI. C, the same gel as in B after staining for in-gel CI activity. D, incorporation efficiency plot was constructed as described in the legend to Fig. 2D. The x axis shows the total amount of tagged subunit in the mitoplasts and the y axis depicts the amount in holo-CI. The solid line (line of identity) indicates maximum incorporation efficiency.
labeling studies provided evidence for the existence of a direct exchange of some CI subunits into CI without the interference of assembly intermediates (8). It was proposed that this mechanism functions to “fix” oxidatively damaged subunits in an efficient manner. However, the relative importance of these two incorporation modes under physiological conditions was never investigated. In this study, we made use of cell lines that conditionally express AcGFP1-tagged CI subunits to investigate the assembly characteristics of different subunits during CI maintenance in a quantitative manner. To this end, we determined the ratio between the amount of tagged subunit present in holo-CI and the total amount of tagged subunit present in mitoplasts at low and high level of steady-state expression, and during the transition between these two states. Of the 14 AcGFP1-tagged CI subunits that were investigated, four (tNDUFS2, tNDUFS7, tNDUFS8, and tNDUFB8) were not or insufficiently incorporated into holo-CI, precluding their further analysis in this study. Importantly, low versus high steady-state expression of the other tagged subunits did not affect CI protein amount and/or activity. This means that none of these subunits is rate-limiting in CI assembly and that subunit-tagging does not detrimentally affect assembly and/or activity of holo-CI.

After 24 h of high level induction, we observed that seven of the tagged subunits displayed a similar fluorescence intensity at the position of holo-CI on a BN-PAGE gel. These subunits included NDUFS1, NDUFV1, and NDUFV2 of the dehydrogenase module, NDUFS3 of the hydrogenase module, and the accessory peripheral arm subunits NDUFS4, NDUFA2, and NDUFA12 of unknown function. Analysis of endogenous subunit levels showed that tNDUFS3 completely outcompeted its endogenous counterpart, indicating that this fluorescence intensity represents the maximum possible level of CI labeling. This demonstrates for the first time that NDUFV1, NDUFV2, NDUFS1, NDUFS3, NDUFS4, NDUFA2, and NDUFA12 are present in a similar stoichiometric amount in mammalian holo-CI. Previous studies have shown that each individual subunit occurs once per CI molecule in the bacterium *Thermus thermophilus* (1, 24), which makes it likely that the same holds true for the much larger mammalian CI. Our data confirm this widely accepted, yet unproven assumption for at least these seven subunits.

Previous studies, which first induced complete depletion of CI by transient inhibition of mitochondrial translation, report CI restoration times of 2–4 days (3, 12–14). The present finding that endogenous NDUFV1, NDUFV2, NDUFS1, NDUFS3,
NDUFS4, NDUFA2, and NDUFA12 were fully replaced by their tagged counterparts in holo-CI after 24 h of induction, indicating that the turnover of these subunits during CI maintenance occurs within this time frame and thus is faster than expected. Indeed, steady-state labeling of holo-CI was reached within 24 h during the transition from low- to high-level expression of tNDUFS1, tNDUFS3, and tNDUFA2. Noticeably, the increase in the amount of tagged subunit in the mitoplasts ceased at the same time that all endogenous NDUFS1, NDUFS3, or NDUFA2 was replaced by its tagged counterpart, indicating that the mitoplast levels of these subunits are tightly regulated. The latter is in concordance with our observation that the holo-CI/mitoplast ratios of the other subunits also remained virtually unchanged under high and low expression levels. The exception was tNDUFS2, which showed maximal incorporation efficiency under leakage conditions but accumulation of monomer subunit on native gel upon induction.

The ability to determine the fraction of total mitoplast fusion protein present in holo-CI revealed that some subunits (tNDUFS1, tNDUFS4, and tNDUFA1) showed maximal incorporation efficiencies at low and high levels of steady-state expression. This strongly suggests that these subunits are directly built in into holo-CI upon import and, in quantitative terms, are not present in assembly intermediates (for schematic representation of our findings, we refer to Fig. 7). In line with this finding, the curves displaying the amount of tNDUFS1 in mitoplasts and holo-CI during the transition from low to high expression levels overlapped. In conclusion, NDUFS1 seems to use predominantly a direct incorporation mode (Fig. 7), compatible with the exchange model (8). In contrast, other subunits had suboptimal incorporation efficiencies. For two subunits with intermediate incorporation efficiencies (tNDUFS3 and tNDUFA2), labeling of mitoplasts clearly preceded that of holo-CI during time induction experiments. In other words, these subunits are primarily incorporated into subassemblies and/or are present as monomers before they are assembled into CI (Fig. 7). Also, tNDUFV1 was found to be present in CI subassemblies. However, in contrast to tNDUFS3 and tNDUFA2, its labeling kinetics of mitoplasts and holo-CI were identical. This strongly suggests that tNDUFV1 makes use of both incorporation modes, in which a certain amount of tNDUFV1 can be directly incorporated into CI upon import, whereas the remainder assemble into smaller (sub)complexes and/or is stored in a NDUFV1 monomer pool (Fig. 7). Newly synthesized subunits were found to assemble at different rates into holo-CI in N. crassa, and it was concluded that this was due to differences in pool size (10). However, our data support the alternative explanation that the different assembly rates reflect the involvement of assembly intermediates.

Obviously, CI biogenesis is much more intricate than represented by the current CI assembly models. For instance, the previous paragraph concluded that two subunits of the dehydrogenase module (tNDUFV1 and tNDUFS1) have distinct incorporation patterns. Several qualitative studies proposed that these two subunits assemble together with other subunits from the same module (3, 4, 25), whereas an in vitro import study reported that NDUFS1 assembled with subunits of the hydrogenase module (8). Both models are very plausible, since NDUFS1 is a functional constituent of both the dehydrogenase and hydrogenase module (24, 26, 27). It is tempting to speculate that NDUFS1 is capable to assemble via both modules, depending on cell type and/or metabolic state. Fig. 1D shows that tNDUFV1 and tNDUFS1 display similar subcomplexes as tNDUFV2 on a BN-PAGE gel probed with an anti-EGFP antibody. NDUFV2 is also a core subunit of the dehydrogenase module (28), suggesting that in our setting these three subunits assemble into dehydrogenase subcomplexes. However, we discussed already that tNDUFV1, in contrast to tNDUFV1 and tNDUFV2, was virtually absent in smaller assemblies after 24 h of induction. The fact that CI labeling for these three subunits was equal, whereas their incorporation levels in subcomplexes differed, makes it unlikely that these subassemblies represent isolation artifacts or breakdown products. Therefore, we propose that these subunits exchange at different rates with already incorporated subunits in subcomplexes during CI homeostasis. This resembles the proposed mechanism for subunit exchange in holo-CI (8).

Recently, it is suggested that assembly intermediates of CI might serve as scaffold for the assembly of other respiratory chain complexes during the formation of respirasomes (14). In line with our findings, the incorporation kinetics of nuclear encoded CI subunits into so called supercomplexes after inhibition of mitochondrial translation occurred for some subunits (NDUFS2 and NDUF9) faster that for others (NDUFS4 and NDUFS1) (14). Our data further suggest that the NDUFV1 pool size is relatively large and has a fast turnover, compatible with the abundance of tNDUFV1 breakdown products. This might be explained by the presence of the NADH binding site, flavin mononucleotide and N3 iron-sulfur cluster (24, 29, 30), which could render this subunit more vulnerable to oxidants than other dehydrogenase subunits.

**FIGURE 7. Proportional use of direct and indirect (delayed) subunit incorporation during CI maintenance.** Incorporation of some subunits into CI occurs immediately after import into the mitochondrion (route A). Other subunits are initially stored in monomer pools and/or assembled into subcomplexes, prior to incorporation into holo-CI (route B). This results in a delay in appearance in holo-CI after import in the mitochondrial matrix. The relative proportion of each incorporation mode (route) for NDUFS1, NDUFS1, NDUFS3, and NDUFA2 is reflected by font size.
Dynamics of Mitochondrial Complex I Assembly

In conclusion, our approach of tagging CI subunits with AcGFP1 provided several new quantitative opportunities to study CI assembly in terms of stoichiometry, incorporation efficiency and temporal assembly dynamics. Subunits of CI are under continuous threat of oxidative damage because CI function is inextricably linked with the production of reactive oxygen species. It is, therefore, important to have detailed knowledge of the mechanisms involved in maintaining the integrity of the complex. This knowledge will contribute to a better understanding of pathologies related to CI dysfunction (31, 32).

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