QST-M-37





Annual Report

202I

Kansai Photon Science Institute Quantum Beam Science Research Directorate National Institutes for Quantum Science and Technology



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はじめに

本年報では、関西光科学研究所(以下、関西研)において 2021 年度に実施された研究開発の主だった 成果を紹介しています。関西研は国立研究開発法人量子科学技術研究開発機構(以下、量研)の研究 開発拠点であり、けいはんな学研都市にある京都府木津地区と兵庫県播磨地区の2か所に研究サイトを持っ ています。そこでは、研究系職員約 80 名及びそれを支える技術系・事務系スタッフを含めて総勢約 150 名の スタッフが、量研における関西研のミッションである「レーザーや放射光による光科学技術の研究開発」を推進し ています。木津地区では、世界トップレベルの高強度レーザー技術を基盤としたレーザー加速やX線発生等の レーザー駆動の新しい放射線源開発、レーザーの短パルス性を活かした超高速計測技術開発、そして放射線 影響や創薬に資する量子生命科学の最先端の研究開発を実施しています。また、播磨地区では、大型放射 光施設 SPring-8 の 2 本の専用ビームラインと計算機シミュレーションを活用することで、新しい放射光 X線利 用技術開発と物質材料科学の最先端研究を展開しています。

量研関西研として6年が経ち、新しい組織の中で優れた研究成果が生まれ始めています。木津地区のレーザ ー科学分野においては、プラズマミラーを用いた J-KAREN-P レーザーのパルスコントラストの改善や、レーザー加 速研究における極薄膜照射実験、次世代のレーザーの基盤技術となる OPCPA 用高安定励起源開発や新し いタイプの発振器開発、そして光と物質の相互作用に関する理論研究などについて着実な成果を挙げました。 量子生命科学分野においては、生物の概日リズム生成に関与するたんぱく質の構造や、発がんに関連したヌク レオソームヒストンの構造メカニズムの解明、そしてモンテカルロシミュレーションを用いた遺伝子突然変異のメカニ ズム解明に向けた研究などを着実に進めています。また、播磨地区においても、放射光 X 線を用いた水素貯蔵 合金開発や物質内部のナノ結晶の観察に加え、反強磁性体におけるスピンダイナミクスの理論に関する研究や、 将来の放射光実験を見据えたデータ取得の機械学習と自動化などの優れた成果を挙げています。さらに、光技 術の社会実装の観点からは、医療応用をめざした小型高輝度光源開発やレーザーによるトンネル検査技術の 実用化に向けた取り組みを引き続き行っています。

関西光科学研究所は、「光」を通じた我が国の量子科学技術の発展とイノベーション戦略に貢献する開かれ た研究拠点としての役割を果たすべく、今後とも職員一同、より一層努力してまいります。皆様のご理解・ご協 力を宜しくお願い申し上げます。



2022 年吉日 関西光科学研究所 所長 河内 哲哉

Preface

This annual report from Kansai Photon Science Institute (KPSI) provides highlights of the scientific and technical research that was conducted over the 2021 fiscal year. KPSI is one of the research and development (R&D) bases of the National Institutes for Quantum Science and Technology (QST) and has two R&D sites, the Kizu site in Keihanna science city in Kyoto prefecture and the Harima site in Hyogo prefecture. There are about 150 staff, comprising around 80 researchers and the technical and administrative staff who support them. We promote the R&D of optical science and technology using lasers and synchrotron-radiation X-rays, which is the mission of KPSI in QST. At the Kizu site, we are conducting state-of-art research such as developing new-type laser-driven radiation sources such as laser accelerated particle beams and ultrashort X-rays based upon world-leading top-class high-intensity laser technology, ultrafast measurement methodology using ultrashort pulse technology, and quantum life science which helps us understand radiation effects and develop new medicines. At the Harima-site, using two contract beamlines of SPring-8 and computer simulation, we are developing new technology to utilize synchrotron radiation X-rays and carrying out state-of-art research in material science.

Six years have passed since the restart of KPSI in QST, and several outstanding results are being produced. In the laser science field at the Kizu site, we have steadily achieved results in areas such as the improvement of the pulse contrast of the J-KAREN-P laser using a plasma mirror, ultra-thin film irradiation experiments in laser acceleration research, development of a highly stable pump source for OPCPA and a new-type oscillator which are expected to become basic technology of next-generation lasers, and theoretical research in laser-matter interaction. In the field of quantum life science, we are steadily proceeding in areas such as the elucidation of the structure of the protein complex involved in the generation of circadian rhythms in living organisms, the establishment of a novel approach for measuring the "Cluster DNA damage" using an atomic force microscope (AFM), and the mechanism of gene mutation using Monte Carlo simulation. Additionally at the Harima site, we have achieved excellent results in areas such as the clarification of the hydrogenation mechanism of Fe-Mo alloy under high pressure, the observation of nanocrystals inside a material via Bragg coherent diffraction imaging using synchrotron X-rays, the theoretical study of spin dynamics in antiferromagnetic materials, and also we have made good progress in machine learning and automating data acquisition which must become important in future synchrotron radiation experiments. Furthermore, from the perspective of social implementation of optical technology, we are continuing efforts to develop compact high-brightness light sources for medical applications and to put laser tunnel inspection technology into practical use.

KPSI will continue to fulfill our role as an open research center of the "science of light" and will contribute to quantum science and technology and the strategy for innovation in Japan. We appreciate your understanding and cooperation.

Good day, 2022 KAWACHI Tetsuya Director General of KPSI

Activities of KPSI

関西光科学研究所の主な動き

【トピックス、シンポジウム・研究会の開催、等】

2021年4月7日から5月27日

【QST 初! クラウドファンディングによる研究開発】

関西研 (木津地区) のジェームズ・ケビン・コーガ博士が 提案した「レーザーで空気中のウイルスを検出できるか?」 について、クラウドファンディングによる研究支援を受け ることができました。2021年4月7日から5月27日の1 か月間での目標金額: 500,000 円に対して、支援総額: 684,029 円を得ることができました。研究成果については、 下記 URL を参照してください。

(日本語) https://academist-cf.com/projects/213?lang=ja (英語) https://academist-cf.com/projects/213?lang=en



クラウドファンディング成功の画面 (アカデミスト Web サイト)

2021年6月22日 (火)

【光・量子ビーム科学合同シンポジウム 2021 (OPTO 2021) 開催】

大阪大学と量研との間で締結された包括協定に基づき、合同シンポジウム(OPTO 2021)を開催致し

ました。年1回開催している合同シンポジウム ですが、昨年に続き今年も会場および Web 配 信によるハイブリッド開催を行いました。参加 者は約 200 名で、大阪大学レーザー科学研究 所と量研関西研を中心に、欧州(ELI-BL)等、 海外からの Web 参加者もおられました。シン ポジウムでは、招待講演・口頭発表、ポスター 発表を中心に若手ポスター賞表彰、大阪大学近 藤賞受賞記念講演、意見交換会が行われまし た。



現地会場での参加者集合写真 2021年6月22日

2021年10月31日(日)

【関西光科学研究所(木津地区) 施設公開】

10月31日に関西研(木津地区)の施設公開を2年ぶりに開催しま した。新型コロナウイルス感染予防対策のため、規模を縮小し、事前予 約制での開催とはなりましたが、100名を超える方々にお越しいただ きました。世界トップクラスの高強度レーザー施設の見学や、光の実 験ショー、工作教室等、ご来場いただいた皆様にはご好評をいただき ました。



光の実験ショー 2021 年 10 月 31 日

2021年12月7日(火)

【セルゲイ ブラノフ上席研究フェロー 外国人叙勲伝 達式 (チェコ共和国)】

功績概要:「科学技術分野における 日本の国際貢献の向 上 に寄与」

令和2年度(2020年度)秋の叙勲において、量研の 上席研究フェローである Sergey Bulanov博士が、外国 人 叙 勲 (Conferment of Decorations on Foreign Nationals)を受章される栄誉に浴されました。新型コ ロナウイルス感染症拡大のために伝達式が1年程度遅 れましたが、無事、チェコ共和国駐箚日本国特命全権大 使 鈴木秀生閣下から褒章を授与されました。



Bulanov 夫妻、鈴木秀生大使夫妻 2021 年 12 月 7 日 在チェコ日本国大使公邸

2022年2月4日(金)

【第5回 理研光量子工学研究センター・量研関西光科学研究所合同セミナー」(Web 開催)】

今回、第5回(5年目)を迎えた本合同セミナーは昨年に続き、完全オンラインで開催されました。参加者は、昨年と同じ程度で常時 40 名以上が参加しました。講演内容は、最先端極短パルス光源開発からレーザー電子加速、分子や物性計測に加え、レーザーを利用した測地やトンネル検査などにおける幅広い領域にわたりました。



左) 講演者リスト



右)Web上での講演者・主催者集合写真 2022年2月4日

2022年3月17日(木)

【日本国駐箚ドイツ連邦共和国特命全権大使 Dr. Clemens von Goetze 閣下来所(視察)】

関西研(木津地区)では、現在ヘルムホルツ研究機構 との学術交流協定をむすび、レーザー加速や高強度場科 学の国際協力を進めています。(ドイツ:ヘルムホルツ 研究機構ドレスデン研究所、日本:大阪大学レーザー 科学研究所および量研関西研)今回、日本国駐箚ドイ



関西研(木津地区)玄関、2022年3月17日(木)

ツ連邦共和国特命全権大使 Dr. Clemens von Goetze 閣下が来所され、研究所の紹介、実験室見学、意見 交換を行いました。今後の研究の進展が期待されています。

2022年3月28日(月)

【QST 未来ラボ次世代放射光利用研究グループ 2021 年度公開ワークショップ】

現在、量研は国の主体として、民間・地域パートナーとともに次世代放射光施設の整備・開発に取り組 んでいます。QST 未来ラボ「次世代放射光利用研究グループ」は、量研が整備する3本の共用ビームラ インを利用して、量子材料や量子生命科学といった量子戦略に関わる研究を行うための基盤を築くこと を目的としています。本ワークショップでは「光電子分光と軟X線磁気分光の融合利用による量子マテ リアルの研究」と題し、第一線で活躍する若手から中堅の4人の研究者による招待講演を含め、現状と 将来展望について活発な議論が行われました。

https://www.qst.go.jp/site/kansai/aslws2021.html

【主な出展等イベント】

7月11日~14日「第21回 SPring-8夏の学校」開催(大型放射光施設 SPring-8、兵庫県佐用町)
11月1日~13日 けいはんな R&D フェア 2021 へ出展(Web 開催)
11月1日~12月1日 けいはんなビジネスメッセ 2021Virtual へ出展(Web 開催)



第 21 回 SPring-8 夏の学校ポスター



SPring-8 の QST ビームライン: BL11XU 表面 X 線回折計での実習の様子

プレスリリース (研究成果)

- 反強磁性モット絶縁物質におけるフェムト秒の電子スピン配列振動を発見

 超高速磁性ダイナミクスの探査手法を提案 (2021年4月26日)
 https://www.qst.go.jp/site/press/20210426.html
 論文タイトル: Antiphase Oscillations in the Time-Resolved Spin Structure Factor of a Photoexcited Mott Insulator
- スクレオソーム中の H2A-H2B テイルの動的構造を解明(2021年7月5日) https://www.qst.go.jp/site/press/20210705.html 論文タイトル: N-terminal tails of histones H2A and H2B adopt two distinct conformations in the nucleosome with contact and reduced contact to DNA
- 3. 希少な元素を使わずにアルミニウムと鉄で水素を蓄える

 水素吸蔵合金開発の新たな展開を先導 (2021 年 7 月 29 日)
 https://www.qst.go.jp/site/press/20210729.html
 論文タイトル: Hydrogen storage by earth-abundant metals, synthesis and characterization of
 - $Al_3FeH_{3.9}$
- 4. レーザー中性子・X 線による同時瞬間撮影
 10万分の1秒の短いパルスで可視光や X 線で見えなかった高速現象を撮る (2021年9月 16日)

https://www.qst.go.jp/site/press/20210916.html

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論文タイトル : Single shot radiography by a bright source of laser driven thermal neutrons and x-rays
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- 5. 巨大負熱膨張のメカニズムを解明
 - さらなる新材料の設計に道を拓く (2021年9月29日)

https://www.qst.go.jp/site/press/20210929.html

論文タイトル: Origin and absence of giant negative thermal expansion in reduced and oxidized Ca_2RuO_4

6. 世界初!元素種を識別して材料のミクロ構造を解析するノイズ耐性の高い新解析法を開発

- 将来的なデバイス材料のミクロ構造研究に活路を開く - (2021 年 12 月 11 日)

https://www.qst.go.jp/site/press/20211211.html

論文タイトル: Bayesian sparse modeling of extended X-ray absorption fine structure to determine interstitial oxygen positions in yttrium oxyhydride epitaxial thin film

- 7. 放射光を使った磁石の奥まで透ける顕微鏡

 X線発光の新原理を用い開発に成功 (2022 年 1 月 25 日)
 https://www.qst.go.jp/site/press/20220125.html
 論文タイトル: Bulk-sensitive magnetic microscope utilizing x-ray magnetic circularly polarized emission
- 8. 世界最強最薄ターゲットによるレーザーイオン加速の実現
 グラフェンと超高強度レーザーが切り拓く極限世界 (2022 年 2 月 16 日)
 https://www.qst.go.jp/site/press/20220216.html
 論文タイトル: Robustness of large-area suspended graphene under interaction with intense laser
- 9. 世界初の完全非破壊コンクリート遠隔打音検査手法を開発
 コンクリート構造物のひび割れをレーザーで素早く視覚化 (2022 年 2 月 22 日)
 https://www.qst.go.jp/site/press/20220222.html
 論文タイトル: Defect detection of concrete in infrastructure based on Rayleigh wave propagation generated by laser-induced plasma shock waves
- 10. 巨大タンパク質複合体による概日リズム制御

- 小角散乱と計算科学の統合アプローチ - (2022年3月10日)

https://www.qst.go.jp/site/press/20220310.html

論文タイトル: Overall structure of fully assembled cyanobacterial KaiABC circadian clock complex by an integrated experimental-computational approach

11. 世界初! 放射線によって生じた DNA 損傷の直接観察に成功

- 老化・がん治療研究にブレイクスルー - (2022 年 3 月 22 日)

https://www.qst.go.jp/site/press/20220322-1.html

論文タイトル: Formation of clustered DNA damage in vivo upon irradiation with ionizing radiation: Visualization and analysis with atomic force microscopy

User Facilities

主要な施設・装置

木津地区

○J-KAREN-P レーザー装置

【装置概要】

世界トップクラスの極短パルス超高強度レーザー です。最大 30 J のレーザーエネルギーを 30 フェムト 秒 (1 フェムトは 1000 兆分の 1) の時間に閉じ込める ことにより 1000 兆ワットの超高強度を実現します。 (右の写真は強力な励起レーザーの光で緑色に光っ ています。)



【供給装置性能】

- ・中心波長:810 nm
- ・繰り返し:シングルショット~0.1 Hz
- ・ターゲット照射エネルギー:~10 J/pulse
- ・パルス幅: 30~50 フェムト秒
- ・集光強度:10²⁰~10²² W/cm²
- ・時間コントラスト比: <10⁻¹¹ (プラズマミラー不使用時) <10⁻¹³ (シングルプラズマミラー使用時)
 - <10 18 (シング)レンノスマミノ 一使用時)
 - <10⁻¹⁵ (ダブルプラズマミラーへアップグレード予定)

上記以外のパラメータもお問い合わせ下さい。

【主要な研究課題】

超高強度レーザー高度化技術開発、イオンや電子の加速技術開発、高輝度 X 線等の量子ビーム源開 発

○QUADRA-T レーザーシステム

【装置概要】

1 秒間に 3000 発のレーザーパルスが繰り出せる高平均出力 ピコ秒パルスレーザーです。

【装置性能】

- ・照射エネルギー: 10 mJ/pulse
- ・波長:1030 nm
- ・繰り返し:3kHz
- ・パルス幅:約1ピコ秒



【主要な研究課題】

高繰り返し高出力レーザー(パラメトリック増幅器等)の開発、高強度テラヘルツ光源の開発

播磨地区

播磨地区では大型放射光施設 SPring-8 に 2 本の QST 専用ビームラインを設置しているほか、日本原 子力研究開発機構(JAEA)の専用ビームラインにも複数の放射光専用実験装置を常設しています。一方 で、QST 専用ビームラインにも、JAEA の専用実験装置が常設されています。

○BL11XU(QST 極限量子ダイナミクス I ビームライン)

【装置概要】

SPring-8標準の真空封止アンジュレータを光源とし、マルチ結晶交換システムを装備することで、 広範囲のエネルギー領域の高輝度放射光 X線を高効率に利用できるビームラインです。

【装置性能】

- ・光源:真空封止アンジュレータ
- ・エネルギー領域:6~70 keV
- ・分光結晶: Si(111)、Si(311)
- ・実験装置:放射光メスバウアー分光装置、共鳴非弾性X線散乱装置、及び表面X線回折計
 - 1. 放射光メスバウアー分光装置

⁵⁷Fe、⁶¹Ni 等のメスバウアー核種を対象とした放射光メスバウアー分光が 可能で、物質の電子、磁気状態から格子振動状態に関する情報などを得るこ とができます。更に、斜入射法や同位体置換試料を利用する事で、金属薄膜 の表面部を原子層単位で測定する事も可能です。

2. 共鳴非弾性 X 線散乱装置

入射X線、散乱(発光)X線の双方のエネルギーを0.1 eV 程度のエネル ギーで分光実験が可能な装置です。4 軸回折計配置で共鳴非弾性X線散乱に より電子励起の運動量依存性が測定できるほか、高エネルギー分解能X線吸 収分光、X線発光分光の実験も可能です。試料温度は10 K から800 K まで 可変です。

3. 表面 X 線回折計

分子線エピタキシー(MBE) チェンバーを搭載した表面構造解析用 X 線 回折計です。半導体量子ドットや半導体多層膜などの成長過程を X 線回折 によりその場観察・リアルタイム観察が可能です。2 台の MBE を交換し、 GaAs、InAs などのヒ素化合物成長と RF-MBE による GaN、InN などの窒 化物半導体成長を行うことができます。



【主要な研究課題】

金属薄膜の原子層単位での磁性探査、遷移金属化合物における電荷・スピン・軌道励起の観測、触 媒や電池電極材料のオペランド電子状態解析、半導体量子ドットや半導体多層膜の成長過程のリアル タイム解析





○BL14B1 (QST 極限量子ダイナミクス II ビームライン)

【装置概要】

偏向電磁石を光源とすることで、連続スペクトルを持つ白色 X 線や高エネルギーの単色 X 線が利用 可能なビームラインです。全反射ミラーや分光結晶の曲げ機構によって、試料位置への集光が可能とな っています。

【装置性能】

- ·光源:偏向電磁石
- ・エネルギー領域: 白色 X 線 (5~150 keV)、単色 X 線 (5~90 keV)
- ・実験装置:高温高圧プレス装置、汎用四軸X線回折計及び分散型XAFS測定装置(JAEA)
- 1. 高温高圧プレス装置

10 GPa(13 万気圧)、2300 K 程度までの圧力・温度状態下の試料を、白色 X 線を用いたエネルギー分散型 X 線回折法やラジオグラフィー法、単色 X 線を用 いた XAFS(X 線吸収微細構造)法や角度分散型 X 線回折法等によって調べる ことができます。

【主要な研究課題】高圧下での金属水素化物形成過程のその場観察

2. 汎用四軸 X 線回折計

令和2年3月にRI実験棟から移設。X線照射位置を可視化するためのレーザ ーを整備しており、がん研究等のための単色 X線を利用した照射実験に使用で きます。高エネルギー単色 X線を利用した回折法により、大型構造材料中の応 力・ひずみ、結晶方位分布計測の測定も可能です。

【主要な研究課題】応力・ひずみ、結晶方位の三次元分布測定、及び照射実験等

○BL22XU(JAEA 専用ビームライン)における放射光専用実験装置

1. ダイヤモンドアンビルセル回折計 (JAEA BL22XU)

最高 70 keV の高エネルギーX 線と大型二次元 X 線検出器の利用により最 大 Q=27Å⁻¹までの X 線全散乱プロファイルが迅速に得られ、約 100Å まで の距離相関の原子二体分布関数が導出できます。専用アタッチメントによ り、室温、1 MPa 未満の水素と窒素ガス雰囲気でのその場観察ができます。

試料-検出器間距離が可変のため高角データから高分解能データまで取得が可能であり、ダイヤモン ドアンビルセルを用いた高圧力下X線回折測定にも適用可能です。

2. 大型 X 線回折計

X線領域のブラッグ反射を利用したコヒーレント X線回折イメージングが実施できる水平振りの大型 X線回折計です。粒径 100 nm 以下のナノ結晶ひと粒のサイズや形状、内部構造(応力、ドメイン等)を含む 3 次元イメージングが可能です。



水素貯蔵合金、負の熱膨張材料、100~500 nm 級チタン酸バリウムナノ結晶粒子一粒の3次元可視 化、40 nm 級パラジウムナノ結晶粒子一粒の3次元可視化

12









施設の稼働実績と利用状況

木津地区

2021 年度の運転時間は J-KAREN-P は 1826 時間、QUADRA-T は 1131 時間、その他装置は合計 1916 時間でした。 J-KAREN-P は上期にコントラスト比を上げるためのプラズマミラー導入作業を行い、下期 より利用研究を再開しました。

実施課題件数

装置名称	独自研究	受託研究	共同研究	施設共用
J-KAREN-P レーザー装置	2	1	0	1
QUADRA-T レーザーシステム	3	1	0	0
kHz チタンサファイアレーザー	6	1	1	0
X線回折装置	1	0	1	0

稼働時間内訳



その他の装置



QUADRA-T



播磨地区

2021 年度の SPring-8 蓄積リングの運転時間は 5208 時間で、放射光利用時間はそのうちの 4440 時間 でした。量研、JAEA とも専用ビームラインでは 10%程度の調整時間を除き、放射光利用時間で独自研 究や受託研究、外部利用者への施設共用と研究支援を行っています。

実施課題件数

ビームライン	独自研究	受託研究	共同研究	施設共用
BL11XU	10	0	3	17
BL14B1	14	0	6	16
BL22XU	6	0	5	3

利用日数

ビームライン	独自研究	受託研究	共同研究	施設共用
BL11XU	95	0	13	56
BL14B1	58	0	21	53
BL22XU	38	0	27	13

*件数、日数ともに QST の利用のみ

*日数は3シフトを1日とした。

稼働時間内訳



Research Highlights

Advanced Photon Research Activities

KONDO Kiminori





The science and technology of advanced lasers are the foundation of the department of Advanced Photon Research. The advanced laser development is the basic part of this department. High-peak-power laser technology, high-repetition-rate and highstability laser technology, and ultrashort-pulse technology are of particular importance. We develop these technologies in-house and apply them to various objects.

The largest laser system in KPSI is the petawatt (PW)-class high-peak-power J-KAREN-P laser. This system has been upgraded since FY2012 by obtaining a supplementary budget. After the long commissioning operation term, internal users in the KPSI and external users started to use this system in the PWclass operation mode. However, from the last fiscal year, we started to set a large plasma mirror system to avoid the rising part of the femtosecond high peak power pulse, which is a serious problem for laser-driven ion acceleration with a nanometer-scale thin foil target. This fiscal year, the machine time of J-KAREN-P for users began in September 2021. Until the middle of September, a large plasma mirror system was installed. Therefore, in the fiscal year 2021, only one theme was shared with external users, although 20 % of all user time was shared in FY2020. Although the user time of J-KAREN-P has declined drastically, the installation of the large plasma mirror system was completed in this fiscal year. Dr. Kon, from the high-intensity laser science group, reports this installation and shows the present performance. Moreover, some ion acceleration experiments, which have been proposed in FY2020 but have not been conducted due to Covid-19, have been conducted in this FY2021. They had the chance to use the extremely focused intensity of

over 10²¹ W/cm² on targets with extremely high contrast, with the suppression of the leading optical component to the main pulse. Unfortunately, for the experimental application of this extreme laser-matter interaction, there is no article in this annual report. Currently, the data analysis is ongoing. We expect to show good results with the large plasma mirror system in the next annual report. This extreme status of our high-power laser infrastructure was maintained by the laser facility operation office, advanced laser group, and high-intensity laser science group. The advanced laser group not only maintained J-KAREN-P to deliver laser pulses with the best conditions but also developed and introduced new technology to maintain the world-leading class condition of J-KAREN-P. Dr. Miyasaka reported the development of a highly stable sub-nanosecond Nd:YAG pump laser, which will improve the synchronization between the pump and seed in OPCPA in the J-KAREN-P laser system. With this improvement, the stability of J-KAREN-P is expected to drastically improve.

The high-intensity laser science group has mainly studied laser-driven ion acceleration, laser electron acceleration, and relativistic high-order harmonic generation with J-KAREN-P in KPSI. One of the most important applications of this technology is the development of laser-driven quantum beams. An extremely high optical field can be formed by focusing an extremely high peak power on an ultimately small spot. The atoms and molecules exposed to this extremely high field were immediately ionized by field ionization. The corresponding optical intensity of the atomic unit was only 3×10^{16} W/cm², which is much lower than that generated with J-KAREN-P. The generated free electrons move along the extremely high optical field, inducing ultra-relativistic



Fig.1 Research and development of the department of Advanced Photon Research

quiver motion. These energetic quivering electrons induced the generation of various secondary radiations. This means that there is the possibility of a compact energetic quantum beam source without conventional accelerator technology. If this technology is established and applied to various fields, destructive innovation can occur. Dr. Fukuda et al., in the advanced laser group, reported energetic ion acceleration with a large-area suspended graphene target. Although the thickness of the graphene target was just a nanometer, this thin foil was irradiated by a high peak intensity J-KAREN-P laser pulse without using a plasma mirror device, and the energetic ions were detected. In the report, they explained what happened in this unique laser shot experiment. Normally, the extremely high peak intensity optical pulse has a highintensity rising edge, which can break away a few-layer graphene thin foil. If so, there should be no chance of accelerating ions with such an extremely thin foil.

Although there are no precise reports in this annual report, one of the most important applications of laser-driven energetic particles is the application to the Quantum Scalpel, which is a new-generation heavy-ion cancer therapy machine. Quantum Scalpel is one of the main projects in QST. The injector part of the quantum scalpel is based on a laser-driven carbon accelerator. The JST-MIRAI R&D program (large-scale type) started in Nov. 2017. This program aims to demonstrate the proof of concept (POC) of a laser-plasma accelerator. In addition to an ion accelerator, a laser-plasma electron accelerator is also under development in KPSI within the MIRAI program. The POC of the laser-driven heavy ion injector must be demonstrated by the end of FY2026. In FY2020, the first-stage gate was successfully passed through, and FY2021 is in the second stage in JST-MIRAI. This year, the driver laser for generating a carbon-ion beam, which is the platform laser for developing the injector of Quantum Scalpel, has been upgraded to a double CPA 30 TW Ti:sapphire laser with a contrast ratio of 10^{10} . With this laser, up to 6th ionized carbon ions were observed, while only up to 4th ionized carbon ions were observed during the1st stage in JST-MIRAI. The beamline for estimating the number of energetic carbon ions with a finite solid angle is estimated to be set in the ion acceleration platform by corroboration with the Sumitomo Heavy Industry. Moreover, using Hitachi Zosen, a repeatable special target system, with which carbon ions are generated, is currently being developed. Highly ionized energetic carbons are expected to be observed at 10 Hz in the near future.

Another important application of advanced laser science and technology is ultrafast probes for life and material science. The ultrafast dynamics group studied not only the basic science of ultrafast molecular behavior, but also developed a unique ultrafast high-power laser system. Until four years ago, the ultrafast dynamics group performed the C-Phost project. In this project, a kHz 10-mJ ps laser system, QUADRA-T, was developed. Recently, this system has been improved using a thin disk amplifier with a higher average power. Moreover, a midinfrared high-average-power laser system is under development for starting attosecond experimental studies in KPSI. These laser systems will be used for laser fabrication and ultrafast laser science, which are studied in the Q-LEAP program at JST. Dr. Otobe reports a semi-classical theoretical approach for lasermetal interaction, which is important in the development of cyber physical system (CPS) laser fabrication in the Q-LEAP project. The ultrafast dynamics group also collaborated with the QST activity in quantum life science. An effective multiphoton microscope is under development for this purpose. In KPSI, a high-repetition-rate mid-infrared laser system with a high average power was developed for this microscope. Dr. Nagashima reported the development of a synchronously pumped OPO with a spatially dispersed beam, which has the potential of an MHz ultrashort high-average-power unique oscillator in the future. A related theoretical study of CPS laser fabrication was also conducted by the X-ray laser group. Dr. Yonetani reported a theoretical account of the calculation of long-range electrostatic interactions and studied on the interaction between DNA and radiation, which is included in the QST activity for quantum life science.

The most important application of our advanced laser science and technology is industrialization. The new SIP program, which follows the previous SIP program that was completed three years ago, began in the last year for the development of nondestructive tunnel inspection technology. This technology is being tested for its application in commercial technology at the venture company Photon-Labo. Non-destructive tunnel inspection is possible using laser hammering technology. This laser hammering technology is expected to be used not only for tunnel inspection abuts but also for various concrete structures. Dr. Hasegawa in the X-ray laser group reports on this study in an attempt to achieve social implementation.

Using high-power laser technology, a strong and compact infrared laser can be developed for various medical applications, including detecting blood glucose measurements. Dr. Akahane reported the development of a compact, high-energy Yb:YAG passive Q-switch laser, which can be used not only for detecting blood glucose measurements but also for pumping nonlinear processes.

Installing plasma mirror system at J-KAREN laser

KON Akira

High-Intensity Laser Science Group, Department of Advanced Photon Research



With the development of chirped pulse amplification technology invented in 1985 [1], research on high-intensity lasers has evolved for various applications [2]. In recent years, laser systems with an output power of more than 10^{20} W/cm² have been constructed in many countries worldwide owing to the development of high-energy and spatio-temporal control technologies by increasing the diameter of optical systems [3].

However, in laser-plasma experiments, target destruction owing to the pre-pulse becomes a problem as the focused intensity increases. The causes of the pre-pulse include amplified spontaneous emission, higher-order dispersion, clipping of the spectrum, scattering from the diffraction gratings, and nonlinear optical effects. These are so intricately intertwined that it is difficult to eliminate their causes.

To solve this problem, plasma mirrors (PMs) have been adopted in experimental facilities to improve temporal contrast [4, 5]. The PM's principle is that the main pulse is reflected by the self-generated plasma on the mirror substrate. However, because of the low intensity of the prepulse, a PM is not created, and most of it passes through the mirror substrate. Also, part of the prepulse is reflected by the anti-reflection (AR) coated mirror substrate surface (Figure.1).



Figure.1 Principle of temporal contrast enhancement using PM

At the Kansai Photon Science Institute (KPSI), Quantum Science and Technology, the upgrade of the J-KAREN laser system and experimental area was completed in 2018 [6]. Also, in 2021 we installed a PM system to improve temporal contrast further for J-KAREN-P (1 PW, 0.1 Hz). Here, we report on the PM system installed at our facility and its performance [7].

As Figure 2 shows, the PM system was installed after the final compressor and before the target chamber. The vacuum chamber's size of the PM system (TOYAMA Co., Ltd.) was $2.1 \text{ m} \times 2.7 \text{ m} \times 2.3 \text{ m}$. The internal components consisted of two periscope pairs, two OAP mirrors, and a PM substrate. The mirror holders of periscopes were the ANTARES series made by LIOP-TEC GmbH, and the driving mechanism was a piezo actuator (Physik Instrumente GmbH & Co. KG). The OAP (TYDEX.

LLC) had a focal length of 2 m (F/8) and a wavefront accuracy of $\lambda/10$ (P-V) at 633 nm. The setup was a "single" PM (see Figure 1). The PM substrate's size was 400 mm × 70 mm × 30 mm, and the laser was incident on the mirror with *S* polarization at an incident angle of 16°. The reflectance of the AR coating on both sides was <1% at 770-830 nm, 16°, and there were silvercoated areas on the top, bottom, and center for alignment. The PM was placed on an automatic stage (Kohzu Precision Co., Ltd) for XYZ and θ x. The mirror was moved for each shot by a drive stage of ±250 mm in the vertical direction and ±35 mm in the horizontal direction. This PM system could be used for ~1,000 shots at 1 Hz without replacing the mirror.



Figure.2 Plasma mirror system at J-KAREN-P



Figure.3 Temporal contrast with and without PM system

Temporal contrast was measured using a third-order crosscorrelator [8] (SEQUOIA, Amplitude Technologies). The performance of the temporal distribution was measured using light picked off from a full beam with a diameter of 25.4 mm. Figure 3 shows the results obtained with and without the PM system. From -10 ps to several hundred femtoseconds, the contrast improved by approximately 100. The trigger time for the PM was expected to be -0.9 ps to -0.3 ps from Figure.3. The dotted line in Figure 3 shows the predicted results without a PM, considering the reflectivity of the PM (65%) and AR coating (1%). The estimated and measured results correlated. However, the temporal contrast could be further improved by one to two orders of magnitude by optimizing the AR coating [9].

The reflectivity of the PM system was calculated using an energy-calibrated charge coupled device after the final stage amplification and an absolutely calibrated integrating sphere and spectrometer after the PM for each shot. In the experiment, reflectance data were obtained when the fluence was varied by changing the distance of the OAP to the PM and laser energy. Figure 4 shows the PM system's reflectivity results. The reflectivity increased as the fluence increased, reaching 85% at a fluence of approximately 100 kJ/cm².

We reported on the configuration and performance of a single PM system installed in the J-KAREN laser system. In future, we plan to conduct experiments using high-contrast lasers (e.g., ion acceleration). Subsequently, we will upgrade to a double PM and adopt a deformable mirror after the compressor to upgrade the laser system.



Figure.4 Reflectivity of PM system

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Energetic ion acceleration by direct irradiation of ultra-intense laser on large-area suspended graphene

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Atomically thin graphene is transparent, highly electrically and thermally conductive, lightweight, and the strongest material. To date, graphene has been used in many applications, including transport, medicine, electronics, energy, defense, and desalination. In this paper, we report another disruptive application of graphene in laser-ion acceleration, in which the unique features of graphene play a crucial role. Osaka University researchers, in collaboration with researchers at the National Institute for Quantum Science and Technology (QST), Kobe University, and National Central University in Taiwan, have demonstrated direct energetic ion acceleration by irradiating the world's thinnest and strongest graphene target using an ultraintense J-KAREN-P laser at the Kansai Photon Science Institute, QST [1].

Recent developments in laser technology have enabled us to access the radiation regime of laser-ion acceleration with relatively thin targets. However, a thinner target is less durable than a thicker target and can be easily broken by a pedestal or prepulse through impact and heating before arriving at the main laser. A possible solution to this problem is to use a plasma mirror, which is a surface plasma produced by the foot of the laser pulse on an optically transparent material that functions as an effective mirror only for the main laser peak. Thus far, diamond-like carbon (DLC) has been used to enhance ion acceleration in an extremely thin target range (< 10 nm) with plasma mirrors. It is necessary to use plasma mirrors, even in a moderately thin target range (10-100 nm), to realize energetic ion generation. However, DLC is not a two-dimensional (2D) material; therefore, it is costly to make it thin and flat. Moreover, graphene is stronger than diamond in an extremely thin region and is more suitable for mass production. Furthermore, installing and operating plasma mirrors at a high repetition rate is expensive. In this study, we investigated another direct solution using graphene as the thinnest and strongest target ever made. We developed a facile transfer method to fabricate large-area suspended graphene (LSG) as a target for laser-ion acceleration with precision down to a single atomic layer [2]. Direct irradiation of LSG targets with an ultraintense laser generates energetic carbons and protons, demonstrating the durability of graphene without a plasma mirror. This application extends the new frontier of graphene science in extreme electromagnetic fields, such as energy frontiers and nuclear fusion.

Experiments were performed using J-KAREN-P at QST-KPSI [3], where the laser intensity reaches the highest class curr-

(a) Laser axis -KAREN Stack 45 LSG Targer normal. NS. (b) 20 2D (c) 3000 **50** μm 1000 1200 1400 1600 1800 2000 2200 Raman (cm⁻¹) 2400 2600 2800

Fig. 1. (a) Schematic of experimental setup with large-area suspended graphene target (LSG). (b) Raman spectrum and (c) optical microscopic images for typical LSG, respectively.

ently available worldwide. Fig. 1(a) shows schematic images of the experimental setup. Fig. 1(b) and (c) show the Raman spectrum and optical microscopic images of a typical LSG. respectively. Here, the narrow bandwidth of the 2D band (< 30 cm^{-1}) and the higher intensity ratio (> 2.5) of the 2D and G bands (I[2D]/I[G]) confirmed the presence of a single-layer LSG. The small peak next to the 2D peak comes from the slight curvature of the LSGlower owing to the high aspect ratio of the LSG. We placed the convex and concave sides of the laser and detector, respectively, to achieve improved beam collimation. The ideal LSG was 0.34 nm, whereas the obtained transferring graphene was close to 1 nm because of the molecular adsorption on the surface [2]. By transferring the graphene layer by layer, we controlled the target thickness with 1 nm accuracy.

The J-KAREN-P laser was focused with an F/1.3 off-axis parabolic mirror on the LSG from 45° to the target normal direction to prevent the back reflection of incident light and destroy the upstream optics without a plasma mirror. The pulse

energy, duration, focal spot, and intensity were ~10 J on the target, 40 fs, ~2 μ m, and ~5 × 10²¹ Wcm⁻², respectively, and measured immediately before the experiment. Energetic ion beams were produced by irradiating the LSGs with intense lasers. The accelerated ions were detected using a stack of radiochromic films, a solid-state nuclear track detector (CR-39), and a Thomson parabola spectrometer (TPS). When a solid target was irradiated with an intense laser, a proton beam was generated independently of the target material, owing to surface contamination from moisture in the air. CR-39s enable us to distinguish heavier ions and mostly carbons from protons based on the size of ion pits [4]. Furthermore, TPS provides the chargeto-mass ratio and the energy spectra.



Fig. 2. (a) Image captured using TPS with eight-layer LSG, where a micro-channel plate and a charged-couple device were used as detectors. The black and green parabolae correspond to carbon and oxygen ions, respectively. (b) Energy distribution functions calculated from (a) for proton and C^{6+} . (c) Carbon pits on first CR-39 covered with a 12 µm aluminum foil with energy of 12–94 MeV. (d) Proton pits on CR-39 behind aluminum foil, two RCFs, and CR-39 corresponding to the energy of 12.2 – 13.2 MeV. (e) Proton energy distribution function obtained using two-layer LSG.

Figure 2 shows the results for an eight-layer LSG, that is, an 8 nm thick target with the full energy of J-KAREN-P at the best focus. In Fig. 2(a), the raw TPS data are represented by blue, black, and red dotted lines corresponding to the protons, carbon ions, and oxygen ions, respectively. Figure 2(b) shows the proton and carbon energy distribution functions $f(E_{P,C})$, where $E_{P,C}$ are the proton and C⁶⁺, respectively, after subtracting the background. Approximately 30% of the oxygen ions were also present. The ion spectra showed bump-on-tail or monoenergetic features in higher-energy tails. When the proton and carbon were accelerated by the same potential field, which occurred in typical laser-ion acceleration, the carbon energy was six times higher than that of the proton because of the difference in the charge-to-mass ratio. The experimental results showed that the proton energy was slightly higher than 1/6 of the carbon energy. Figure 2(c) and (d)

show the etched pits of mostly carbons and protons, respectively, which are consistent with the TPS. Figure 2(e) shows the same plot as Fig. 2(b), except for the double-layer LSG corresponding to a thickness of 2 nm. This image was captured under a different condition in which the laser incident angle was 10° with a moderate contrast level of ~ 10^{-7} . For the eight-layer LSG shot, the contrast level was ~ 10^{-10} or lower

As mentioned above, the pre-pulse and pedestal are significant practical problems in the extremely thin target region. Recent experiments have utilized single- or double-plasma mirrors to suppress the pre-pulse and pedestal and achieve energetic ion acceleration. Surprisingly, energetic ions are generated by irradiating single-figure-nanometer thin targets with an ultra-intense laser at ~ 10^{22} Wcm⁻² without a plasma mirror. Our double-layer LSG is the thinnest target that generates energetic ions, even without a plasma mirror at this thickness.

In conclusion, the results of this study apply to the development of compact and efficient laser-driven ion accelerators for cancer therapy, laser nuclear fusion, high-energy physics, and laboratory astrophysics. The direct acceleration of energetic ions without a plasma mirror demonstrates the robustness of the LSG. We will use atomic-thin LSG as a target mount to accelerate other materials that cannot stand without support. We also demonstrated energetic ion acceleration at non-relativistic intensity (not discussed here). This research will help us investigate laser-ion acceleration using relatively small laser facilities. Furthermore, energetic ion acceleration was achieved even without a plasma mirror in the extremely thin target region, thus, opening a new region for laser-driven ion acceleration.

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Sub-nanosecond Nd:YAG pump laser for optically synchronized optical parametric chirped-pulse amplification

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The J-KAREN-P laser, which is the petawatt-class femtosecond laser system developed in the Kansai Photon Science Institute (KPSI), achieves a high pulse contrast of 10¹² by combining optical parametric chirped-pulse amplification (OPCPA) as a pre-amplifier and subsequent Ti:sapphire amplifications [1]. In the current J-KAREN-P laser, the timing jitter between the pump and seed pulses for the OPCPA is ~100 ps, because the two oscillators for the pump and seed lasers are electrically synchronized with a commercially available pulse delay generator. Reducing the timing jitter improves the stability of the output pulse conditions of the J-KAREN-P laser to achieve higher-quality experiments. Teisset et al. developed an optical synchronization scheme with a single oscillator by generating pump pulses from a portion of the signal pulses to reduce timing jitter [2]. A timing jitter of less than 400 fs is realized by optically synchronized pumping; however, the pulse duration reported to date is too short for our purpose. To use the optically synchronized pump pulses for the OPCPA in the J-KAREN-P laser, a long pulse duration is required because the pulse duration of the signal pulses is stretched to ~1ns to avoid optical damage in the subsequent Ti:sapphire amplifiers. A frequency-doubled Nd:YAG laser was used for OPCPA pumping in the J-KAREN-P laser because of the efficient laser operation at room temperature without complex cooling systems. The longest pulse duration of an optically synchronized Nd:YAG laser was reported to be 207 ps [3]. Herein, we report the development of a highly stable optically synchronized frequency-doubled Nd:YAG pump laser. Pulse durations of 600 ps at 1064 nm and 490 ps at 532 nm were obtained by optimizing the Nd:YAG crystal temperature [4].

Figure 1 shows the schematic of the proposed optically synchronized pump laser. All components were set on a watercooled optical table to minimize the temperature effects. The laser comprised a Ti:sapphire oscillator, fiber systems for stable chirped-seed pulse generation at a wavelength of 1064 nm, bulk amplifier systems for high-energy amplification, and a frequency doubler. Pulses from a Ti:sapphire oscillator were focused on a 23 cm photonic crystal fiber to extend the spectrum to include 1064 nm. Polarizing the input pulses was controlled by a feedback control system composed of a spectrometer and half-wave plate mounted on a motorized rotation stage to stabilize the output spectrum at 1064 nm. The stabilized pulses were amplified to 130 mW (rms: 0.2 %) by four Yb-doped fiber amplifiers and







Fig. 2. Measured pulse shapes before passing through the

stretched to 1 ns by a chirped Bragg fiber grating. The repetition rate of the fiber output pulses was reduced from 80 MHz to 10 Hz using a Pockels cell for subsequent amplifications with a diode-pumped Nd:YAG regenerative amplifier and a two-pass main amplifier. The regenerative amplifier was designed to set a beam diameter of ~2 mm on a 6 mm diameter Nd:YAG rod to avoid Fresnel diffraction from the rod edge. With 10 round trips, the pulses were amplified to 20 mJ (RMS: 0.3%) with a beam diameter of 2.3 mm. The amplified pulses were expanded to a diameter of 4 mm before injecting into the main amplifier. A serrated aperture of 5 mm diameter was placed after the beam expander to avoid optical damage induced by Frenel diffraction at the 8 mm diameter Nd:YAG rod in the main amplifier. A pulse energy of 200 mJ was achieved with excellent stability of better than 0.2% (rms). The amplified pulses' frequency was doubled in an LBO crystal with a length of 13 mm. A pulse energy of 130 mJ with a stability of 0.6% (rms) at a wavelength of 532 nm was obtained, with a conversion efficiency of 64%. The pulse duration after frequency doubling was shortened to 330 ps; whereas, the pulse duration before regenerative amplification was 1 ns. Fig. 2 shows the measured pulse shapes before and after the regenerative amplifier with a 30 GHz oscilloscope. The pulse duration after applying the regenerative amplifier was 360 ps. The pulse duration decreased as the number of round trips in the regenerative amplifier increased.We investigated the phenomenon of pulse shortening owing to the pulse shape distortion behavior of high-gain chirped-pulse amplification in a regenerative amplifier. Assuming a single value of stimulated emission cross-section for chirped pulse amplification in a regenerative amplifier, a pulse duration of more than 850 ps was calculated using the Frantz-Nodvik equation [5]. The difference between the experimental and calculation results was derived from the wavelength dependence of the stimulated emission



Fig. 3. Measured (solid line) and calculated (dashed line) pulse shapes after the regenerative amplifier.

cross-section [6]. The temperature dependence and narrow bandwidth of the stimulated emission cross-section of the Nd:YAG crystal affected pulse shortening. In this experiment, the Nd:YAG crystal was cooled with water at 18°C. The dashed line in Fig. 3 shows the calculated pulse shape after regenerative amplification at a rod temperature of 31.5°C. The pulse duration was increased to 620 ps because the wavelength of the peak cross section shifted to the spectrum peak of the chirped pulse. To confirm this effect experimentally, we measured the amplification at a cooling-water temperature of 31°C. The solid line in Fig. 3 shows the pulse shape after regenerative amplification. The pulse duration was extended to 600 ps, which agreed with the calculation results. After the main amplification, the pulse duration was reduced to 540 ps. Therefore, we considered the pulse shortening to be due to the gain narrowing caused by the decrease in stored energy when the latter part of the pulse was amplified. The pulse duration after doubling was 490 ps. A longer pulse duration will be achieved using a chirped fiber Bragg grating with a higher group velocity dispersion and a lower bandwidth. The increase in the chirp rate of the incident pulse enabled the regenerative amplifier to generate a longer pulse for the optically synchronized OPCPA in the J-KAREN-P laser.

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Synchronously pumped optical parametric oscillator with a spatially dispersed beam

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Synchronously pumped optical parametric oscillator (SPOPO) is a conventional apparatus used for wavelength-tunable infrared light sources [1–3]. Periodically poled materials have been used in several SPOPO experiments because these materials provide several advantages such as high nonlinear coefficients and lack of spatial walk-off [4-6]. In addition, they have a significant advantage that various structures with different poling periods can be flexibly designed [7]. Furthermore, structures with chirped or fan-out poling periods have been developed [8-10].

Chen et al. proposed a method for an optical parametric amplifier (OPA) using a fan-out type poled device [11]. In this method, broadband phase matching was achieved by adjusting the fan-out poling period and the spatially dispersed beam. The frequency-domain OPA is a similar method that uses a spatially dispersed beam [12]. In this study, the demonstration of a broadband SPOPO is presented using a fan-out type poled device and a spatially dispersed beam [13].



Fig. 1. Optical layout of the SPOPO. It consists of components such as spherical lens (SL), cylindrical lens (CL), periodically poled LiTaO₃ (PPLT), dichroic mirrors (DM1 and DM2) that reflect only the signal light, and an output coupler (OC) with a signal light transmittance of 50%. The inset depicts the output signal beam profile measured at a distance of 1.7 m from the OC.

An Yb-doped fiber laser was used as the pump source for the SPOPO. The laser had a repetition frequency of 43.3 MHz, central wavelength of 1038 nm, full-width at half-maximum (FWHM) pulse length of 390 fs, and a maximum pump power of 14 W. Fig. 1 shows the optical layout of SPOPO. This layout was similar to that used in a previous study [14]. MgO-doped periodically poled LiTaO₃ (MgO:PPLT) was used as the nonlinear material. The output coupler (OC) had a constant transmission rate of 50% in the frequency range of 1300 – 1700 nm. The cavity length was 3.46 m, which corresponded to the repetition frequency of the pump laser. Flat mirrors were used in the cavity for dichroic mirrors (DM1 and DM2) and the OC. Only one spherical lens was used in the cavity. The spherical lens was made of CaF₂ with a broadband anti-reflection (AR) coating and it had a focal length of 254.7 mm for a wavelength of 1580 nm.

An N-SF11 glass equilateral prism was installed in the cavity. Fig. 2 shows the details of the fan-out type MgO:PPLT and the spatially dispersed beam. The prism had an AR coating for the wavelength range of 1500 - 1700 nm when the S-polarization light was incident at an angle of 60.6° . The prism generated an angular dispersion of $\Delta\theta/\Delta\lambda$. The spatial displacement at the spherical lens was $\Delta X/\Delta \lambda = f\Delta \theta/\Delta \lambda = 9.3 \times 10^{-3}$ mm/nm, where f is the focal length. In Fig. 2, the three lines represent signal beam axes with different wavelengths. The beams were parallel between DM1 and the spherical lens, and coaxially overlapped between the prism and OC.



Fig. 2. Details of the fan-out type MgO:PPLT and the spatially dispersed beam.

The nonlinear crystal was 1 mol% MgO-doped stoichiometric PPLT (manufactured by OXIDE). The crystal had a broadband AR coating in the range of 1000 – 1700 nm and it had a length (Z-direction), height, and width (X-direction) of 10, 2, and 6 mm, respectively. The effective width was 4.5 mm, where the poling period had a nonlinear fan-out structure and it varied in the range of 28.6 – 32.4 μ m. Fig. 3(a) shows the relationship between the position X in the MgO:PPLT and the poling period. The fan-out structure was designed to exhibit a linear relationship between horizontal position (X) and the signal wavelength, as shown in Fig. 3(b). The slope of the linear relation was $\Delta X/\Delta \lambda = 9.24 \times 10^{-3}$ mm/nm, which was equal to the value obtained by the prism dispersion.



Fig. 3. (a) Poling period of MgO:PPLT as a function of the horizontal position (X). (b) Signal wavelength that satisfies the phase-matching condition as a function of X.

The spectral dispersion in the cavity was dominated by the dispersion of the transmission materials (MgO:PPLT, CaF₂, and N-SF11 glass). The thickness of the CaF2 lens was 2.7 mm and the pass length in the N-SF11 prism was approximately 8 mm. The GDD values were 567, -3.7, and, 246 fs² for MgO:PPLT, CaF₂, and N-SF11, respectively. Therefore, the total GDD was 809 fs². This spectral dispersion was compensated for by using a fused silica (SiO₂) plate with an AR coating. Fig. 4(a) shows the

group indices of the transmission materials $(n_g(\lambda))$. The plotted values represent the difference from the value at $\lambda_0 = 1580$ nm. The total dispersion was evaluated as a summation of group indices multiplied by their lengths, that is, $\Sigma \{n_g(\lambda) - n_g(\lambda_0)\}L$, where L is the length. The length of the SiO₂ plate was determined to be 24 mm to compensate for the total dispersion. Fig. 4(b) depicts the total dispersions with and without the SiO₂ plate.



Fig. 4. (a) Group indices of transmission materials in the SPOPO cavity as a function of the wavelength. (b) Total dispersion of transmission materials in the SPOPO cavity.

The pump beam was linearly focused in the horizontal direction using a cylindrical lens with a focal length of 200 mm. A wide region in the MgO:PPLT was pumped and broad spectral components of the signal light were generated in this line-focus setup.



Fig. 5. (a) Signal output power as a function of pump power when the OC position is zero. (b) Contour plot of the signal spectra as a function of the OC position when the pump power is 14.0 W.

Fig. 5(a) depicts signal output power as a function of the pump power. The OC position was determined to maximize the signal output power and this position was defined as the origin of the OC position. The threshold pump power was approximately 7 W. The slope efficiency was approximately 20% in the pump power range over 10 W. Fig. 5(b) shows a contour plot of the spectra as a function of the OC position when the pump power was 14.0 W. The figure shows that the spectra expand with an increase in the OC position. The spectral width was approximately 80 nm at the OC position of 0.0 and 120 nm at the OC position of 5.0 μ m. The broadband oscillation was observed in a wider range of the OC position with an increase in the pump power.

The pump pulse length (FWHM) was measured as 392 fs using the SHG-FROG (frequency-resolved optical gating) measurement. The signal pulse length was 81 fs from the SHG-FROG for the line-focus setup with a pump power of 14 W and the OC position at zero. This length was approximately 1/5 of the pump pulse length. It was observed that the signal pulse length was the shortest when the OC position was near zero. Furthermore, when the OC position was over zero, the spectrum became broader, but the pulse length did not decrease.

In conclusion, the broadband operation of SPOPO with a spatially dispersed beam was demonstrated. Additionally, signal pulses with a pulse length of approximately 1/5 of the pump pulse length were generated.

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Semi-classical approach for laser-metal interaction

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The comprehensive modeling of laser material machining [1-3] is complex, with multi-scale time and space, multi-phase, and possibly chemical reactions. For laser processing of metals, a two-temperature model (TTM) has been proposed as the simplest continuum model [4]. In the TTM, empirical parameters are employed for all material properties, and the penetration depth models the dynamics of the electromagnetic field. The most critical assumption of the TTM is the quasi-equilibrium of the electron temperature and the electron-lattice interaction term.

Recently, the first-principles approach employing timedependent density functional theory (TDDFT) [5] has been applied to laser-matter interactions [6,7]. Although TDDFT offers a compromise between accuracy and computational feasibility, its computational cost remains high. Electron-electron collisions (which are not included in TDDFT) play an important role in the laser-metal interactions.

The Vlasov equation has been employed in nuclear physics and electron dynamics in metal clusters to describe the collision process of fermi particles [8-10]. Because the Vlasov equation treats the distribution function of space, momentum, and time, it requires tremendous computational resources. In general, the computational cost is reduced by assuming that the distribution function is the summary quasi-particles.

The time-dependent Kohn–Sham equation [5] is the fundamental equation of the TDDFT.

$$i\hbar\frac{\partial\phi_i}{\partial t} = H_{KS}\phi_i \qquad 1$$

Here:

$$H_{KS}[\mathbf{n}_{e}(\vec{\mathbf{r}}, \mathbf{t})] = -\frac{\hbar^{2}}{2m}\nabla^{2} + V_{eff}[\mathbf{n}_{e}(\vec{\mathbf{r}}, \mathbf{t})]$$

where H_{KS} was the Kohn–Sham Hamiltonian, *m* the electron mass, V_{eff} the effective potential, and $n_e(\vec{r}, t)$ the time-dependent electron density.

$$n_{e}(\vec{r},t) = \sum_{i=1}^{N} |\phi_{i}(\vec{r},t)|^{2}$$
3

Further, the von Neumann equation governed the time-evolution of the density matrix $\hat{\rho}$

д

$$\frac{\hat{\rho}(\vec{r},\vec{r}',t)}{\partial t} = -\frac{i}{\hbar} \Big[\hat{H}_{KS}, \hat{\rho}(\vec{r},\vec{r}',t) \Big].$$

$$4$$

By performing the Wigner transformation and taking the limit $\hbar \rightarrow 0$, the density operator was mapped onto a real function $f(\vec{r}, \vec{p}, t)$, which obeyed the Vlasov equation:

 $\frac{\partial}{\partial t}f(\vec{r},\vec{p},t) = -\frac{\vec{p}}{m} \cdot \nabla_{\vec{r}}f(\vec{r},\vec{p},t) + \nabla_{\vec{r}}V_{eff} \cdot \nabla_{\vec{p}}f(\vec{r},\vec{p},t), \quad 5$ Here, $f(\vec{r},\vec{p},t)$ was interpreted as the electron distribution function in phase space.

The effective potential was a function of the electron

density $n_e(\vec{r},t)$ and decomposed into

 $V_{eff}[n_e(\vec{r},t)] = V_{Cou}[n_e(\vec{r},t)] + V_{xc}[n_e(\vec{r},t)] + V_{ext}(\vec{r},t)$, 6 with the exchange-correlation potential V_{xc} [11], external field potential V_{ext} , and

$$V_{Cou}[n_{e}(\vec{r},t)] = \sum_{j} V_{ps}(\vec{r}-\vec{r}'_{j}) + V_{H}[n_{e}(\vec{r},t)],$$
 7

where V_{ps} , V_H , and j denoted the ionic pseudopotential, Hartree potential, and label of ions, respectively.

We employed a modified Heine–Abarenkov-type local pseudopotential for V_{ps} with parameters to reproduce the allelectron density functional theory calculations [12]. V_H was evaluated by solving the Poisson equation. V_{xc} was the exchange-correlation potential in the local density approximation. The laser-electron interaction was described in the length gauge.

We approximated the six-dimensional time-dependent function with the summation of the pseudo particle (PP):

$$f(\vec{r}, \vec{p}, t) = \frac{1}{N_s} \sum_{k=1}^{N_{pp}} g_r \big(\vec{r} - \vec{r}_k(t) \big) g_p \big(\vec{p} - \vec{p}_k(t) \big)$$
8

Here, k was the label of the PP, and N_{pp} the total number of PPs defined as $N_{pp} = N_s N_e$, where N_s and N_e were the number of PPs per electron and the total number of electrons, respectively.



Fig. 1 (a) Conductivity and (b) refractive indexes of bulk-Al with the Vlasov equation and TDDFT [15]. The filled circles indicate the experimental results [13].



Fig. 2 Laser fluence dependence of the energy absorption with a wavelength of (a) 800 nm, (b) 1,030 nm, and (c) 1,200 nm.[15]

We defined the smoothing functions for position $g_r(\vec{r})$ and momentum $g_p(\vec{p})$ in the Gaussian form. The Newton equation described the motion of the PP under an effective potential with a periodic boundary condition as follows:

$$\frac{d\vec{r}_k}{dt} = \frac{\vec{p}_k}{m}$$

$$\frac{dp_k}{dt} = -\int d\vec{r} \, V_{eff}(\vec{r}) \nabla_{\vec{r}} g_r(\vec{r}_k(t) - \vec{r}) \tag{10}$$

The dynamics of PPs were performed by the Verlet method [14].

We discussed the complex optical conductivity, refractive index, extinction coefficient, and reflectivity as functions of photon energy [15]. Despite the simplicity of the Vlasov approach, the results agreed with the TDDFT results and experimental values (Fig. 1). The peak and dip at approximately 1.5 eV in the TDDFT results were due to inter-band transition, which was not reproduced by the present Vlasov approach because the latter considered only the free-electron dispersion.

Fig. 2 shows the absorbed energy as a function of laser intensity with laser wavelengths of (a) 800 nm, (b) 1,030 nm, and (c) 1,200 nm [15]. The triangles indicated the results obtained using TDDFT, and the circles indicated the results obtained using

the Vlasov equation. The scaled one-photon absorption was indicated by the solid line for the guiding eye. At all wavelengths, the Vlasov equation agreed with TDDFT below the intensity of 1×10^{13} W/cm². Above 1×10^{13} W/cm², Vlasov underestimated compared with TDDFT.

Here, we proposed a semi-classical approach for the lasermetal interactions employing the Vlasov equation. The master equation and potential were based on the Kohn–Sham Hamiltonian with a local density approximation. We employed PP to reproduce the continuous density distribution function in the real and momentum spaces. Our results agreed with TDDFT for the weak and intense laser field regimes. In particular, the Vlasov equation reproduced the optical properties of aluminum without artificial parameters. The advantages of the Vlasov equation was its low computational cost (approximately 1/50 of TDDFT) and high scalability. Assuming Pauli blocking, we included the electron-electron collision process in the materials. Further, the collision between PP and the pseudo-potential ensured semiclassical molecular dynamics.

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Social implementation of laser hammering system for maintenance of concrete structures

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1. Introduction

In Japanese road tunnels, periodic inspections are required once every five years in the wake of the collapse of a ceiling panel in the Sasago Tunnel on the Chuo Expressway in December 2012. Maintenance works are carried out in accordance with the inspection guidelines established by the Ministry of Land, Infrastructure, Transport and Tourism, but there are concerns about a shortage of inspectors in the near future due to aging and retirement. Therefore, new technologies for measurement/ monitoring and non-destructive inspection are being registered in the "Inspection Support Technology Performance Catalog" led by the Ministry of Land, Infrastructure, Transport and Tourism. QST and Photon Lab Co., Ltd. improved the inspection speed of the "laser hammering method"¹) that involves non-contact inspection and digitization. This project was collaboration work with the Institute for Laser Technology (ILT) which was supported through the SIP: Cross-ministerial Strategic Innovation Promotion Program (2014-2019, Yozo Fujino PD) led by the Cabinet Office in collaboration. Based on this result, we have developed a new laser hammering system²⁾ assuming social implementation. In June 2020, this system was registered in the Inspection Support Technology Performance Catalog (registration number: TN020003-V0020)³⁾, and trial work was started with a construction consultant company that carries out inspection works⁴⁻⁷⁾.

2. Laser hammering system

The main requirements for measurement/monitoring and infrastructure inspection technologies are improvement of the inspection speed, remote control, quantification, and digitization, which are in good agreement with the characteristics of the laser hammering method. A schematic of the laser hammering method is shown in Figure 1(a). In this method, a high-intensity pulse laser (1064 nm, 10 Hz, Nd:YAG laser) was used as the impact force, instead of a hammer. The irradiated laser energy was converted into momentum as a blowout of small particles (laser ablation), and an impact force was generated on the concrete surface. The excited vibration on the concrete surface was measured using an interferometer (laser Doppler vibrometer) with a continuous oscillation laser (532 nm, Nd:YVO4 laser). A wavelength of 532 nm is suitable for the surface vibration of concrete because it passes through water and can measure surface vibrations even if the concrete surface is wet. Both lasers were irradiated to the inspection point through galvanic mirrors for high-speed scanning. The interval between each measurement point was set arbitrarily (minimum 10 mm), and the range of one scan was approximately 500 mm for the object at a distance of 5 m. If a 500 mm square area is measured at 50 mm scan intervals, 121 points are measured, and the required time is 36 s (measured three times per point to improve the S/N ratio).

A schematic for detecting defects using the laser hammering method is shown in Figure 1(b). Resonant vibration occurs in the area containing a defect, such as peeling inside the concrete, due to interference in the elastic wave at the position of the defect (deflection resonance). For example, in the case of a disk-shaped defect with diameter 2a located at a depth l, the resonance frequency is proportional to l/a^2 . On the other hand, resonant vibration does not appear in sound (defect-free) areas. Inspectors using the conventional hammering test recognize the difference in the magnitude and frequency of these vibrations as a difference in sound, and determine the soundness of the concrete. Similarly, in the laser hammering method, the detection of defects is determined by the resonant frequency and spectral power.



Figure 1) Scheme of the laser hammering method⁴⁾

Figure 2 shows the laser hammering system (LHS) mounted on a 4-ton loading platform truck. The LHS can be easily transported to the destination tunnel, and laser alignment after arriving at the tunnel is almost unnecessary. The LHS contains an optical system (impact laser, laser vibrometer, scanning, and aiming system), a control system (laser driver, chiller, device for data acquisition and analysis), and a diesel electric generator. Each component was mounted on vibration isolators to prevent interference due to vibrations from the diesel electric generator and vehicle engine. Sub-micron scale vibration measurement is possible when the vehicle is stopped⁵).



Figure 2) Laser hammering system mounted on vehicle⁴⁾

The impact and detection laser beams are guided to the inspection points, such as cracked areas of the tunnel surface and locations where the presence of defects was reported in previous inspections, by the aiming mirror installed at the laser output port. By using the aiming mirror, it is possible to measure any position in the circumferential direction of the tunnel, and several meters in the travel direction of the tunnel, while the vehicle is stopped. For measurements of more than several meters in the travel direction, the vehicle is moved.

3. Application to maintenance of road tunnels

A trial to apply LHS to the maintenance of road tunnels was started in collaboration with a construction consultant company in 2020⁴⁻⁷). Figure 3 shows inspection of a horizontal construction joint (spring line), which are required to be inspected according to Road Tunnel Periodic Inspection Guidelines. This work was carried out by placing an LHS vehicle in a lane closed to traffic. The area around the LHS was designated as a laser-controlled area according to JIS C6802 "Safety of laser products".



Figure 3) Laser hammering inspection under one-sided traffic restrictions in road tunnel⁴⁾

Figure 4 shows an example of the results for one span of the spring line (10.5 m) and the measured vibration spectrum. The inspection area per scan was 50×50 cm, and the inspection interval was 5 cm. The areas shown in red were identified by LHS as containing defects, and the green areas were determined to be defect-free. The defects were concentrated near the boundary of the spring line, and this result was in good agreement with that obtained by a conventional hammering test.

The results in Figure 4 result in a binary judgment (sound or defective). However, it is possible to obtain more detailed information by analyzing the vibration spectrum. For example, the relative severity of a defect can be determined from the spectral power ((2) >> (4) > (1) >> (3)). Furthermore, by recording the vibration spectrum, it is possible to obtain information on the progress of the severity of the defect in subsequent inspections.

It will be possible to develop AI that can classify the soundness according to categories I to IV in response to the requirements of the Road Tunnel Periodic Inspection Guidelines by learning the vibration spectrum and time waveform of various defects. This research is in progress as the improvement of the judgment/diagnosis system of LHS in the SIP 2nd phase (2018~, Yuichiro Anzai PD).

4. Summary

The laser hammering system developed by QST and Photon Lab has been used in several trials⁴⁻⁷) as a support tool for the inspection of road tunnels since June 2020. To meet the demands of practical usage, such as increased speed and smaller size, improvements and optimization⁸) of the device and operation are in progress through multiple national projects, including the SIP 2nd phase.

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Figure 4) Inspection result at the horizontal construction joint (spring line) in the road tunnel⁴)

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Theoretical account on the calculation of long-range electrostatic interactions

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Electrostatic interactions are critically important because they govern the microscopic and macroscopic behaviors of various molecular systems. The calculation of electrostatic interactions has been one of the major subjects in particle simulations, such as molecular dynamics (MD) and Monte Carlo (MC) simulations [1, 2]. Short-range interactions, such as van der Waals forces, can be evaluated using a cutoff treatment that considers finite-range contribution only. However, electrostatic interactions are longrange, and the contributions from distant regions are nonnegligible. Thus, a simple cutoff treatment suffers from various artifacts in terms of the structural and energetic properties. The most significant problem is that the cutoff treatment does not ensure convergence of the simulation results. Increasing the cutoff length does not necessarily improve the simulation results. A typical example is the structure of liquid water; previous MD studies [3-5] clearly showed that increasing the cutoff length worsens the result, contrary to our expectations. In extreme situations, the cutoff simulation produces an artificial layer structure in which water molecules are highly ordered in dipolar orientations [3-5].

In 1980, Neumann and Steinhauser [6] explored the theoretical background of the cutoff problem. Using the Onsager dielectric continuum model with a cutoff boundary condition, they evaluated the electric behavior of a dielectric medium surrounding a dipolar particle. The result of the distancedependent Kirkwood function $G_{K}(R)$ showed that the cutoff boundary condition produced field polarization that was strikingly different from the correct no-cutoff case. A no-cutoff calculation that considers the contributions from an infinite region provides a flat $G_{K}(R)$ profile. However, when the cutoff condition was applied, a peculiar hole-like profile appeared in the $G_{K}(R)$ function. This emergence of unusual polarization can be recognized as cutoff-induced dipolar ordering. In this way, Neumann and Steinhauser [6] theoretically demonstrated how the cutoff treatment produces an artificial orientational order in dipolar systems.

Recently, cutoff-based real-space treatment has improved significantly [7–9]. Suitable tuning of shifting and damping can substantially reduce cutoff artifacts. MD simulations using this approach have provided good results for various molecular systems, such as liquid water, liquid–vapor, biomolecules, and inorganic materials. However, such developments in real-space treatment have not yet been completed. Studies on the real-space treatments are trying to address the fundamental problem, that is, if long-range electrostatic features in condensed phases can be described using the short-range treatment. To achieve this,



Fig. 1: Dielectric continuum model (left) and the twodimensional view (right). The parameters a and r_c are the radii of the central dipolar particle and cutoff, respectively. b denotes the region of the dielectric continuum. (Reprinted from [10], with the permission of AIP publishing)

thorough theoretical exploration is required.

In this study [10], we employed the dielectric continuum model (Fig. 1) to explore the fundamental background of the recently developed shift treatments. Shifting has been widely used as a pivotal part of modern real-space treatment. From a physical point of view, shifting can be interpreted as an image-charge (or image multipole) effect that accounts for the electric cancellation in condensed phases [7]. Here, we selected three ways of shifting: shift potential (SP), gradient-shift force (GSF), and Taylor-shift force (TSF) [9]. We explored the physical background of these shifting schemes using the Neumann-Steinhauaer approach [6]. The results clarified how shifting reduces the cutoff artifact and what remains to be solved [9].

Fig. 2 shows the results of the theoretical analysis. When using a simple cutoff (i.e., bare), $G_k(R)$ decreased when $R < r_c$ and then increased when $R > r_c$. This is a *hole-shaped artifact*, known as a major cutoff artifact [3–6]. In contrast, SP, GSF, and TSF exhibited distinct behavior, in which $G_k(R)$ gradually increased with R. For comparison, the no-cutoff result [6] is also shown in the figure. In this case, $G_k(R) = 1$ for any value of R. All other results show some deviations from this correct result. SP, GSF, and TSF avoided the cutoff artifact of the hole shape. However, deviations from the flat profile [i.e., $G_k(R) = 1$] remained, indicating that some artificial orientational order existed. That is, shifting alone was not sufficient for SP, GSF, and TSF; shifting should be employed with damping, as suggested previously [9].

To clarify why different treatments produce different shapes of $G_K(R)$, we next considered the relationship between the dipolar


Fig. 2: Distance-dependent Kirkwood function $G_K(R)$. The parameters used were a = 1.0, $r_c = 4.0$ and b = 6.0, which are denoted by the vertical dotted lines. No-cutoff [6] and reaction field (RF) results overlapped completely. (Reprinted from [10], with the permission of AIP publishing)

force *K* and $G_K(R)$. Based on the dielectric continuum model (Fig. 1) employed here, the central dipolar force *K* produces the surrounding dielectric polarization *P*, which consequently produces the electric field *F*. The spatial distribution of the *F* vectors seemed similar between the bare cutoff and the shift treatments; however, we observed a significant difference. The difference from the no-cutoff result $\Delta F (= F - F_{no-cutoff})$ clearly shows that the bare cutoff was distinctly different from the other cases: SP, GSF, and TSF. In particular, the ΔF of the bare cutoff was discontinuous at the cutoff boundary $r = r_c$, where the directions were upset. Such drastic changes were not observed in SP, GSF, and TSF.

To verify the theoretical results, we performed Monte Carlo simulations on dipolar particles [$r_c = 9.38$, 17.31, and 28.00 Å]. The MC results (Fig. 6 in Ref. [10]) were slightly different from the continuum results (Fig. 2), particularly at the short cutoff $r_c = 9.38$ and 17.31 Å. However, at the largest cutoff of $r_c = 28.00$ Å, the MC result was slightly similar to the result of the continuum model. The continuum feature was more visible when the number of particles included within the spherical cutoff region $r = r_c$ increased (that is, $r_c \sim 28.00$ Å). This indicates that increasing the particle number made the local microscopic feature less dominant and the global macroscopic feature more dominant. In

other words, the continuum model analysis captured the feature with a large r_c well.

In particular, the continuum analysis presented here could predict if *hole-shaped artifact* appeared; it produced *hole-shaped artifact* for bare cutoff, but not for other shift treatments (Fig. 2). This indicates that the shift treatments resolved the $G_K(R)$ problem of the simple cutoff. In this respect, the present continuum model results were consistent with the MC results.

As shown in the MC results (Fig. 6 in Ref. [10]), the bare cutoff exhibited a large deviation, which became significant as r_c increased. At $r_c = 28.00$ Å, the hole shape grew significantly. In this case, an artificial layer structure [3–5] was formed (Fig. 3; bare cutoff). The other shift treatments did not show such an artificial structure (Fig. 3). This suggests that the shift treatments still resolved this problem. This observation was consistent with that for the continuum model, where shift treatments did not show any indication, that is, no hole.

This study [10] theoretically explains how recent modifications to electrostatic calculations can improve the simple cutoff. The present study, using the continuum model, offers a stringent criterion to determine whether the basic electrostatic behavior is correctly described. This knowledge will be useful for future construction of electrostatic treatments. In addition, the present study contributes to the progress of computational physics, which covers a broad range of scientific fields, such as material and biological sciences.

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Fig. 3: Results from the Monte Carlo simulations of dipolar particles. Each particle is colored red and gray to show the dipolar direction. Bare cutoff shows the layer structure (denoted by the yellow dashed lines), whereas GSF and no-cutoff (Ewald) do not. (Reprinted from [10], with the permission of AIP publishing)

Development of compact, high-energy Yb:YAG passive Q-switch laser for pumping nonlinear process

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We studied tiny ytterbium (Yb)-doped high-intense passive Qswitched lasers for the development of palm-sized intense infrared (IR) lasers that employ optical-parametric oscillation (OPO). The passive Q-switched laser can generate intense, short laser pulses without opto-electric polarizing devices such as Pockels cell; it is suitable for pumping palm-sized OPO lasers. When generating mid-IR (over 3 µm) laser pulses with an OPO scheme, the low OPO gain and low laser damage threshold (<10 MW/cm²) of nonlinear OPO crystals should be considered. A laser cavity for this mid-IR OPO must have a relatively long (a few centimeters) OPO crystal and be pumped by long nanosecond (ns: 10⁻⁹s; a few ns ~10 ns) laser pulses. Yb:YAG was chosen as laser gain media to generate such long ns pulses. For the same cavity length, shown in Fig.1, considerably longer ns pules can be generated with Yb:YAG laser cavity than with conventional Nd:YAG¹. In this study, we have designed compact laser cavity with Yb:YAG gain media, and succeeded to generate such long laser pulses from passive Q-switched pump lasers with high energy.



Fig. 1 Calculated width of output laser pulse with respect to optical cavity length in a passive Q-switch laser with a Cr:YAG saturable absorber. For the same cavity length, considerably longer pules can be generated from an Yb:YAG laser cavity (green) than Nd:YAG (pink).

The schematic of our passive Q-switch laser is shown in Fig. 2. Yb:YAG ceramics is used as gain medium, which is monolithically calcined with Cr:YAG ceramic material as a saturable absorber. The Yb:YAG-side surface of the monolithic ceramic was high-reflection (HR) coated, which acted as a cavity mirror. The composite structure of Yb:YAG/Cr:YAG monolithic ceramics has the advantage of reducing the number of boundary surfaces for higher damage resistance. The Yb:YAG ceramic was pumped by fiber-coupled laser diode (LD) at 940 nm. The laser cavity of Yb:YAG microchip laser comprised the HR coated surface of Yb:YAG ceramic and partially reflecting coated surface of the output coupler. The optical length of the laser cavity was approximately 24 mm. The Q-switched pulses generated at a wavelength of 1030 nm, as shown on the left picture in Fig.2, are launched from the surface of the output coupler. In the experiment,



Fig. 2 Schematic and photo of Yb:YAG microchip laser. LD: laser diode, FO: focusing optics, HR: high-reflection coating, OC: output coupler.

laser pulses of 3 mJ, 3 ns (FWHM) were obtained at a repetition rate of 60 Hz.

We also measured the long-term stability of laser output of the microchip laser. As seen in Fig. 3, the energy of the output laser pulses fluctuated because of the effect of residual thermal load on the pump LD irradiation. In the first experiment, the standard deviation of the measured output laser energy was 12% with water cooling. To improve stability of the laser, thermoelectric (TE) cooling was introduced with the laser system. After this modification, the standard deviation was reduced to 5%, which is sufficient stability for pumping a mid-IR OPO.

A plural Q-switch laser pulse generation from a single-pump LD irradiation is available with this laser cavity. When the duration of LD pump pulse exceeds the next Q-switching threshold, successive Q-switchings occur in a sequence. In the typical temporal waveform shown in Fig.4, we can see a barrage of four Q-switched laser pulses on a single LD pump pulse irradiation. There are several merits with plural Q-switching operation on OPO pumping, such as repetition multiplexing, high power and



Fig. 3 Long-term energy output of microchip laser. Standard deviation of the output laser energy was 12% with a water-cooled jacket (red), which was improved to 5% with a thermoelectric cooled jacket (blue).



Fig. 4 (Top) Observed plural passive Q-switching; typical temporal waveform of output laser pulses. (Bottom) total output energy and laser efficiency with respect to pump LD pulsewidth. Inset numbers (1-4) indicate the numbers of plural Q-switching pulses.

efficiency of laser output, which is a potential tools for compact mid-IR lasers.

The laser cavity could be extended generate a longer (~10 ns) energetic laser pulse. Figure 5 shows the measured output energies and pulse durations with respect to the cavity length of the Yb:YAG microchip laser. As the theoretical prediction², pulse duration of output laser pulse is almost proportional to cavity length, and lengthening the cavity increases the output laser pulse energy with cavity mode expansion. The output energy, shown in Fig. 5, increases with a longer cavity, and a maximum 6.3 mJ pulse energy was measured with a cavity of ~160 mm. The measured pulse duration was also increased with a longer cavity from 4.3 ns to 12.0 ns (FWHM).



Fig. 5 Output energies (red circles) and pulse durations (blue triangles) with respect to cavity length in the Yb:YAG passive Q-switch laser, which is increased with cavity lengthening. Maximum energy and pulse duration were 6.3 mJ and 12.0 ns (FWHM), respectively, with a \sim 160 mm cavity.

Spatial and temporal shaping of laser pulse should be considered to utilize high-intensity passive Q-switch laser pulses as the OPO pump pulse. In general, laser pulses generated in long-cavity lasers tend to exhibit multimodal beam profiles. In a long cavity passive Q-switch laser using a saturable absorber, a laser pulse that originates from the large-diameter multimode part contains a low intensity temporal component generated from the outer higher-modes of the laser cavity. The low-intensity temporal component does not contribute to nonlinear laser interaction, such as OPO in some cases, which has a threshold of laser intensity. And furthermore, irradiation by the un-needed lowintensity part of the laser pulse causes laser damage on the OPO crystal. Spatial filtering (SF) was introduced to eliminate the lowintensity, higher-mode of Q-switched output laser pulses. As shown in Fig. 6, a spatial filter, comprising a lens pair and a pinhole was installed on the output laser optical path. The diameter of the pinhole was set to ~1.2 times larger than the diffraction-limited size of the laser beam, which is a compromise



Fig. 6 (Top) Schematic of spatial filtering in Yb:YAG passive Qswitch laser. (Bottom) Spatial profiles of output laser pulses: left: without spatial filter, right: with spatial filter.

between eliminating higher modes and maintaining the transmission efficiency.

After SF, shown in lower images in Fig.6, the spatial profile was significantly improved. The duration of the laser pulse was also reduced from 11.4 ns to 10.8 ns with SF, and the threshold pump laser intensity for some mid-IR OPO was reduced from 3.7 MW/cm² to 2.3 MW/cm², which clearly shows the effectiveness of the SF technic.

In summary, we developed a compact, intense passive Q-switch laser with Yb:YAG/Cr:YAG composite ceramics for pumping nonlinear processes. In the microchip laser cavity of 24 mm length, an output laser pulse with 3 mJ of energy and 3 ns of duration was obtained at 60 Hz. We also observed multiple Qswitched laser oscillation on a single LD pumping, which is expected to efficient laser pulse generation. In a long cavity design, the energy and the pulse duration of the generated Qswitched laser pulse were increased to 6.3 mJ and 12.0 ns, respectively, which can be a more powerful infrared OPO pump source with spatial filtering. Lasers studied in this research will contribute to the development of palm-sized intense mid-infrared lasers, which have many applications in fields including medical care, environment measuring, material treatment.

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Research Activities at the Synchrotron Radiation Research Center

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A wide range of X-ray techniques are being developed at the Synchrotron Radiation Research Center (SRRC) for the investigation of the structural, electronic, and magnetic properties of matter, primarily using the two QST beamlines, BL11XU and BL14B1, at the large synchrotron radiation facility SPring-8. Hard X-ray non-destructive *in situ* techniques are investigated to enable ground-breaking measurements using techniques such as single atomic-layer magnetic microscopy, element/orbitalspecific excited state analysis, coherent X-ray nanoscale imaging, and time-resolved studies of samples under extreme conditions.

The SRRC consists of the Coherent X-ray Research Group, High-Pressure Science and Stress Research Group, Magnetism Research Group, Condensed Matter Theory Group, and a beamline operation office (Table 1). The research activities of each research group are summarized below.

Measures have been undertaken to support digital transformation (DX) of the activities at SRCC by utilizing the supplemental budget to upgrade the equipment and facilitate automatic measurements. A contract was signed with the Advanced Research Infrastructure for Materials and Nanotechnology in Japan (ARIM) to initiate a project for the development of a unifying structure for the data generated by the in-house analysis equipment at the facility. Additionally, work has been conducted to upgrade the monochromator of BL11XU for the reduction of beam instabilities and enhancement of the beam quality. The equipment developed at SRCC is available for public use and we accepted 34 research proposals in FY2021 as a member institute of the Nanotechnology Platform Japan [1].

Synchrotron/ free-electron laser (FEL) X-ray facilities are being constructed worldwide at a rapid pace. Research is being conducted by the Coherent X-ray Research Group to ensure state-of-the-art synchrotron/FEL research. Advanced measurement/analysis techniques are being developed for the effective utilization of novel advanced light sources. The application of spatially coherent X-rays has enabled the visualization of the inhomogeneity inside matter. An apparatus for Bragg coherent X-ray diffraction imaging (Bragg-CDI) is being developed and it will be applied to the study of inhomogeneous structures of sub-micrometer-sized crystalline fine particles of functional materials. The technique is applicable to particles with a size in the range of 40 - 500 nm [2].

Nonlinear and quantum optical techniques are being developed using simple atomic systems to exploit the temporal coherence and intensity offered by next-generation light sources at short wavelengths. Previously, the soft X-ray beamline of SACLA was used to observe 'superfluorescence' at extreme ultraviolet wavelengths. A source of superfluid liquid helium droplets is being developed for use at SACLA. The development of this source will extend the technique to shorter wavelengths and ensure improved pulse control. The development and application of world-leading pump-probe [3] and innovative [4] gas-phase measurements are ensured by this group.

A surface X-ray diffractometer coupled with a molecular beam epitaxy chamber was used for in situ measurements of crystal truncation rod (CTR) scattering to investigate the atomic structure of gallium nitride surfaces under growth conditions. The presence of a pseudo 1×1 structure was confirmed by the CTR profile analysis and the structural parameters (interlayer distance, coverage, and temperature factor) were determined.

The research objectives of the *High-Pressure Science and Stress Research Group* are (i) the development of experimental techniques for in-situ measurements under extreme conditions, including high-pressure conditions and compressed hydrogen gas environments (ii) fundamental and applied studies of advanced functional materials using the above-mentioned techniques (iii) studies on synchrotron radiation X-ray irradiation effects on tumors. The research group members focus on the high-pressure synthesis of novel hydrogen-rich compounds and the study of the nano- to meso-scale structures of functional materials by collaboration with the Coherent X-ray Research Group.

Metal-hydrogen systems are a primary research target of this group. A hydride Al₃FeH₄ at 9 GPa and 750 °C was synthesized [5]. In-situ X-ray and neutron diffraction experiments clarified the hydrogenation mechanism of Fe-Mo alloy under high pressure [6]. An atomic pair-distribution function (PDF) is obtained by Fourier transformation of the total scattering pattern and it is a powerful tool for studying the local structures of functional materials. The nitrogen gas flow type temperature control system was improved to enable measurement between 100 - 400 K. This system was applied to the giant negative thermal expansion oxide Ca2RuO4 to elucidate the mechanism of negative thermal expansion [7]. Additionally, studies have been conducted to investigate the irradiation effects on tumors with nanoparticles containing high-Z elements. The ablation of the tumor spheroid with iodine-loaded nanoparticles was observed using X-rays (E = 33.2 keV) above the iodine Kabsorption edge [8].

The spin degrees of freedom of electrons have a significant effect on functional materials and devices, ranging from high- T_c superconducting oxides to spintronic devices. In the *Magnetism Research Group*, advanced X-ray spectroscopic techniques such as nuclear resonant scattering (NRS), resonant inelastic X-ray scattering (RIXS), and X-ray magnetic circularly polarized emission (XMCPE), and measurement techniques utilizing measurement informatics are being developed to unveil the functionality of such materials and devices. Measurement informatics aims to improve the efficiency of measurement and analysis using machine learning techniques. Recently, active learning is applied to spectral measurements to enhance its experimental efficiency [9]. Automated stopping criterion based on expected generalization error is revealed to be effective in Xray absorption spectroscopy experiment with active learning. This method enables high-throughput autonomous experiments for the discovery of novel materials. For NRS, synchrotron-based Mössbauer spectroscopy using a frequency-domain technique was conducted by incorporating a nuclear Bragg monochromator. The microscopic origin of the interfacial perpendicular magnetic anisotropy (PMA) on the Fe/Au(111) multilayer surface was studied by 57Fe probe layer method. The measured spectra exhibited the PMA characteristics and enhancement of the interfacial component in the hyperfine field through Fe-Au bonding. The results demonstrated that the PMA in the Fe/Au(111) interface originated from the interaction between the spin and orbital magnetic moments in Fe and Rashba-type spinorbit coupling in Au [10]. XMCPE is a novel magnetic spectroscopy technique operated in the hard X-ray regime with a distinctive feature of a large flipping ratio (25%) for the $K\alpha$ emission of 3d transition metal elements. Recently, a bulksensitive magnetic microscope was developed utilizing XMCPE with a 10-µm lateral resolution and the observation of magnetic domains in an electrical steel sheet was demonstrated [11]. Additionally, the sum rules of XMCPE at L edges for 3dtransition metal elements were deduced utilizing the theoretical framework developed for Ka XMCPE spectra [12].

The *Condensed Matter Theory Group* develops advanced simulation methods based on quantum mechanics to investigate the theory of condensed matter using supercomputers. Additionally, the methods are applied to perform numerical simulations to understand various properties of materials such as

magnetism, high-Tc superconductivity, and catalytic activity, with the support of X-ray experiments such as from SPring-8. The present activities of this group include (i) the development of advanced simulation techniques based on first principles path integral molecular dynamics for investigating the nuclear quantum effects of hydrogen-containing materials such as clathrate hydrates (ii) the development of a numerically exact method diagonalization to investigate time-resolved spectroscopies of antiferromagnetic Mott insulators, which can provide insights into their electronic dynamics [13] (iii) the development of a theoretical framework and computational codes to analyze material properties probed by X-ray spectroscopies such as X-ray magnetic circular dichroism (XMCD), RIXS, and XMCPE [12], and (iv) theoretical investigations using large-scale numerical exact diagonalization of quantum spin systems found in hypermaterials, which exhibit novel quantum phases such as spin nematic liquid.

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Members	K. Ohwada (GL), J. Harries, T. Sasaki	H. Saitoh (GL), A. Machida, A. Shiro, N. Oshime	T. Inami (GL), T. Mitsui, H. Iwasawa, T. Ueno, K. Fujiwara, A. Koide, (K. Ishii, A. Agui)	(T. Sakai (GL)), T. Ikeda, K. Tsutsui, T. Nomura
Typical techniques & Apparatuses	Coherent x-ray scattering Surface x-ray diffraction Quantum optics in EUV region	High-pressure and high- temperature x-ray diffraction Pair distribution function analysis	Mössbauer spectroscopy Resonant inelastic x-ray scattering X-ray magnetic circularly polarized emission	First principles molecular dynamics simulations, numerical calculations
Research objective	Ferroelectrics, nitride semiconductors, superfluorescence	hydrogen containing materials, negative thermal expansion materials, irradiation effects on tumor	magnetism, spintronics, high- <i>T_c</i> superconductivity, measurement informatics	hydrogen containing materials, high- <i>T_c</i> superconductivity, catalysis, magnets
Beamline Operation Office/ Y. Katayama(Section Manager), K. Sugawara, Y. Teraoka, Y. Kai, S. Fujikawa, A. Shimada				

Table 1. Groups at the Synchrotron Radiation Research Center.

Hydrogen storage by earth-abundant metals, high-pressure synthesis of Al₃FeH₄

SAITOH Hiroyuki



High Pressure Science and Stress Research Group, Synchrotron Radiation Research Center

Hydrogen is a promising energy carrier because it does not emit greenhouse gases during usage. Additionally, once the energy is converted to hydrogen, it will not be lost even if it is stored for a long period of time. Hydrogen is a gas at ambient conditions, and therefore, has a low volumetric energy density. In addition, the high risk of explosions and high diffusivity in the material, which causes a decrease in the material strength called hydrogen embrittlement, make the storage of hydrogen even more difficult. Along with high-pressure hydrogen gas tanks, liquid hydrogen, chemical hydrides, and hydrogen storage materials have been extensively studied for a long time.

In hydrogen storage materials, hydrogen is stored in the form of atoms. A schematic of hydrogen storage in a hydrogen storage material is shown in Fig. 1. Molecular hydrogen dissociates on the alloy surface, diffuses into the alloy as atomic hydrogen, and occupies specific sites in the metal lattice to form interstitial hydrides. As a result, hydrogen exists in a solid state, and the volumetric density of hydrogen in hydrogen storage materials is comparable to that of liquid hydrogen, despite the presence of metal atoms that make up the hydrogen storage materials.

As aforedescribed, hydrogen storage materials have excellent hydrogen storage properties, but still certain problems must be solved, such as low gravimetric hydrogen density and high material costs. Among the hydrogen storage materials that have been developed thus far, LaNi₅ and TiFe alloys are commercially used; however, LaNi₅ contains an expensive and heavy rare earth element, and TiFe contains relatively expensive titanium. Is it possible to obtain a new hydrogen storage alloy that can solve these issues through material exploration?

The hydrogen storage alloys reported thus far are composed of a combination of metals with high and low affinity to hydrogen [1]. The guideline of "combining metals with high and low affinity for hydrogen" for the synthesis of hydrogen storage materials suggests that hydrides of metals with high affinity for hydrogen are extremely stable to release hydrogen; therefore, the addition of metals with low affinity for hydrogen is adjusted to release hydrogen at moderate temperature conditions. Hence, it is difficult to obtain hydrogen storage materials from only metals



Figure 1. Schematic of hydrogenation reaction of a hydrogen storage material.

with a low affinity for hydrogen, and there are no actual reports of such materials. Many studies have been conducted on the search for materials based on conventional guidelines, and the author believes that it will be difficult to realize new classes of materials.

We have been searching for new classes of hydrides consisting only of metals with a low affinity for hydrogen using the high-pressure hydrogenation technique, which is expected to be advantageous for the synthesis of new hydrides. Aluminum is a lightweight and inexpensive metal that is ideal for use as a raw material for hydrogen storage materials. However, it is also known that aluminum and its alloys have very low affinity for hydrogen. In 2021, we reported that an alloy consisting only of aluminum and iron, earth-abundant metals with low hydrogen affinity, can form hydrides at high temperatures and pressures and that the resulting hydrides may be thermodynamically stable even at near-ambient pressures [2]. If the synthesis pressure of this hydride can be lowered, it can be used as a hydrogen storage material. The experimental procedure and results are described below.

The starting material was an $Al_{13}Fe_4$ alloy prepared in an arc furnace. The ingot obtained was crushed into a disk with a diameter of 1 mm and height of 0.4 mm. The disk-shaped sample was set in a sample capsule made of boron nitride and located in a cubic pressure medium along with a graphite heater, internal hydrogen source, and hydrogen-sealing capsule. The internal hydrogen source evolved hydrogen at a high pressure, and the evolved hydrogen was confined in the hydrogen-sealing capsule. Hydrogen can permeate the boron nitride capsule and react with the sample, whereas byproducts from the internal hydrogen source cannot pass through the capsule. Thus, we hydrogenated the alloys under a high pressure and temperature.

The sample was pressurized to 9 GPa at room temperature and heated to 750 °C at approximately constant pressure of 9 GPa. The sample was immersed in fluid hydrogen to form its hydride. The sample was then cooled to room temperature and depressurized to ambient pressure. The hydrogenation process of the sample was monitored by an in situ synchrotron radiation Xray diffraction measurement system installed on beamline BL14B1, SPring-8 [3].

Figure 2 shows in situ synchrotron radiation X-ray diffraction profiles of the sample under high pressure and temperature. The bottom profile was taken when the sample was pressurized to 9 GPa at room temperature. All the Bragg peaks that appeared in the profile were indexed by a unit cell of $Al_{13}Fe_4$ alloy, indicating that no structural phase transition occurred during pressurization. When the sample was heated to around 600 °C, the internal hydrogen source evolved hydrogen, and the sample was immersed in fluid hydrogen. Simultaneous to the hydrogen evolution, the peaks intensities from $Al_{13}Fe_4$ began decreasing and new Bragg peaks began to appear. The Bragg peaks from $Al_{13}Fe_4$ vanished approximately 5 min after the



Figure 2. Series of in situ synchrotron radiation X-ray diffraction profiles recorded when Al₁₃Fe₄ alloy was hydrogenated to 9 GPa.

sample was heated to 750 °C. The newly appeared Bragg peaks were indexed by a unit cell of a novel hydride, Al₃FeH₄. Thus, we determined the hydrogenation pressure–temperature conditions of Al₁₃Fe₄ alloy under high pressure and temperature using in situ synchrotron radiation X-ray diffraction technique. The obtained Al₃FeH₄ retained its crystal structure during the process of cooling to room temperature and decompression to ambient pressure and could be recovered at ambient conditions.

The obtained hydride recovered at ambient conditions was examined for hydrogen release upon heating using a thermogravimetric-mass spectrometer. The hydrogen evolution from the hydride started at approximately 150 °C. The gravimetric hydrogen density calculated from the weight loss during hydrogen release was 2.9 wt%. This value is comparable to that of typical hydrogen storage alloys such as LaNi₅ (1.4 wt%) and TiFe (1.9 wt%). The crystal structure of the sample after hydrogen release was the same as that of Al₁₃Fe4.

Because the decomposition temperature and pressure conditions of hydrides can be determined using in situ synchrotron radiation powder X-ray diffraction, the thermodynamic stability of hydrides can be evaluated from these results. The results showed that Al₃FeH₄ had the potential to be thermodynamically stable near ambient conditions; that is, thermodynamically, this alloy had the potential to be used for hydrogen storage near ambient pressure. At present, however, the hydrogenation reaction requires a high pressure of more than 7 GPa. We believe that this is because the chemically stable surface oxide layer on the Al₁₃Fe₄ alloy inhibits the hydrogenation reaction. If the effect of the oxide layer could be suppressed in some way, hydrogen storage could be realized using only aluminum and iron, which are the cheapest and most industrially important metal combinations. We are currently studying the reduction in the hydrogenation pressure.

Why does Al₃FeH₄ remain stable at ambient conditions even though it deviates from the conventional guidelines for hydrogen storage materials? We believe that the key point lies in the bonding between metal and hydrogen. To investigate the stabilization mechanism of Al₃FeH₄, the crystal structure was investigated using neutron and synchrotron radiation X-ray diffraction techniques. A schematic of the crystal structure obtained is shown in Fig. 3. The arrangement of the metal atoms



Figure 3. Schematic of the crystal structure of Al₃FeH₄.

changed significantly owing to hydrogenation. This indicates that Al₃FeH₄ may be a different type of hydride from the interstitial hydride, in which hydrogen is located at interstitial sites in the metal lattice.

The bond length between aluminum and hydrogen and between iron and hydrogen in Al3FeH4 were compared with those in known hydrides, and the results showed that they were close to those in hydrides with a covalent nature. It was also observed that the structural units in Al₃FeH₄ were similar, but not perfectly matched, to the typical structural units found in hydrides with a covalent nature. For example, observing the bonds between Fe and hydrogen in Al₃FeH₄, the bond lengths and arrangements were noted to be similar to those of octahedral [FeH₆]⁴⁻ complex anions present in other iron complex hydrides. However, it was also observed that the structural unit in Al₃FeH₄ had 1/3 partial occupancy of six hydrogen sites and was distorted from the octahedral structure. It is likely that hydrides that deviate from conventional guidelines are stabilized by a peculiar bonding nature that does not fit the conventional classification of hydrides. In the future, we aim to obtain guidelines for the synthesis of this new class of hydrides through further experimental elucidation of their electronic structures.

Acknowledgments

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Development of an apparatus for Bragg coherent X-ray diffraction imaging at SPring-8

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The demand for structural characterization in nano- to micrometer scales, also called mesoscale regions, is rapidly increasing. For example, in the field of ferroelectrics, the size effect [1-3] -- the dielectric properties of ferroelectric fine particles strongly depend on the particle size -- is well-known but their structural origin is undiscovered. In the case of BaTiO₃ fine particles, which have attracted considerable scientific and industrial interest over the past few decades, the dielectric constants have been reported to increase as the average particle size decreases and rapidly decrease below the critical size [4,5]. This cannot be confirmed even when we consider only perovskite-type unit cells. Currently, it is suggested that the particle is composed of a core region with a tetragonal crystal system and a shell region with a cubic crystal system. A gradient lattice strain layer (GLSL) [6] bridges the core and shell regions, and nanosized ferroelectric domains exist in the core region. To obtain a true picture of what occurs within a BaTiO₃ fine particle, it is essential to precisely investigate the structure of individual single particles, including the particle size, shape, and global structure (such as domains and GLSL) in nano- to-micrometer scales.

Coherent X-ray diffraction imaging (CDI) [7] is a highly effective technique for visualizing the structure of nano- to micrometer particles. CDI is a lens-less imaging technique that was rapidly developed following the availability of coherent Xray light sources, the 3rd generation synchrotron light sources, such as SPring-8. Several CDI techniques are in use at present [8]; however, the following approach is common to all techniques. If an object is illuminated by coherent X-rays, the far-field diffracted wave is proportional to its Fourier transform. This situation guarantees the feasibility of image reconstruction. However, the diffracted images are actually measured as a square of the Fourier transform, and the phase information of the diffracted wavefield is lost; therefore, we cannot obtain the image using a simple inverse Fourier transform. This is called a phase problem. However, the image of an object can be reconstructed using a phase-retrieval algorithm. The diffracted images must be sampled at a frequency that is finer than the Nyquist interval.

Here, we focus on the Bragg CDI [9] technique for visualizing individual fine particles. Bragg-CDI is a method dedicated to crystalline fine particles, in which Bragg scattering can be obtained. In Bragg-CDI, a fine particle is illuminated by coherent X-rays and the diffraction patterns around the Bragg position are measured in three dimensions. Therefore, a three-dimensional image of the fine particles can be obtained by the phase retrieval of the diffraction patterns. The retrieved amplitude and phase are related to the electron density and displacement/strain field of the crystal lattice in the fine particles, respectively [10]. Because the Bragg scattering profile and intensity are sensitive to the degree of decoherence of the crystal lattice, the Bragg CDI method can sensitively image the displacement or strain field in fine particles.

Because the orientation of particles is random in a powder, polycrystalline, or ceramic sample, the Bragg-CDI technique is

an efficient tool for investigating individual particles in such samples. Because the Bragg peaks of individual particles can be separated in the scattering space, individual particles can be distinguished from their neighbors. Moreover, in most cases, three-dimensional data can be obtained by changing the crystal angle by an order of 1°, which is an advantage over other CDI techniques.

The Bragg-CDI method and related techniques have already been developed in the United States (Advanced Photon Source) and applied to the study of various nanoparticles [11-14]; however, no dedicated apparatus for Bragg-CDI exists in Japan. Thus, we aim to construct an apparatus for investigating the structure of individual fine particles using the Bragg-CDI method. The Bragg-CDI method and precise X-ray structural analysis of individual fine particles are key techniques for obtaining a true picture of what occurs in a single particle.

In the field of fine particle development, the scanning electron microscope (SEM) is frequently used for characterizing the particles. In this case, the shape and surface of the particles can be observed; however, information on the internal structure (strain and domains) and the shape and surface of the back of the particle cannot be obtained. Furthermore, particles in devices such as deep inside multilayer ceramic capacitors (MLCCs) cannot be measured using electron beams. The use of (hard) Xrays overcomes this disadvantage.

In this study, we develop an apparatus for Bragg-CDI and perform the three-dimensional imaging of some fine particles using the apparatus [15-16].



Fig. 1 Schematic drawing of the setup for Bragg coherent Xray diffraction imaging. Speckle patterns were drawn using ImageJ [17].



Fig. 2 (a) Reconstructed 3D image of a BaTiO₃ fine particle and cross-sectional drawings of norm (corresponds to the electron density) and phase (corresponds to the strain). (b) Phase (strain) distribution for a series of samples: Pd, BT-Y2c, BT-S3, and BT-S5. These figures were drawn using VESTA 3. [19])

: The Bragg-CDI apparatus was developed at BL22XU in SPring-8. Figure 1 shows the schematic of the experimental setup. The X-ray energy is typically 8 keV. Fine particles were dispersed on the SiN membrane or STEM grid, which did not disturb the coherency of the X-rays. The membrane/grid was set on a sample holder and inserted into a vacuum chamber specially designed for Bragg CDI measurements [16]. The X-rays were focused using a refractive lens (Karlsruhe Institute of Technology) to approximately 2×2 (V × H) μ m², with a focal length of 1.0 m. A fine particle on the SiN membrane was illuminated by the focused X-rays, and the diffracted X-rays were collected by a single photon-counting two-dimensional X-ray detector, that is, Eiger500k (Paul Scherrer Institute) or Pilatus300k (DECTRIS). A typical coherent X-ray diffraction pattern is shown in Fig.1. One can clearly see a typical coherent X-ray diffraction pattern, the so-called speckle pattern. We collected three-dimensional speckle patterns around the Bragg position by changing the Bragg angle (ω -scan) to within $\pm 1.0^{\circ}$. with a step size of typically 0.01° . Using the three-dimensional Bragg diffraction data, we reconstructed three dimensional images of the fine particles. Phase retrieval of the observed diffraction patterns was carried out using a series of hybrid input output (HIO) and error reduction (ER) algorithms [18], which are frequently used in CDI. Three types of BaTiO₃ and Pd particles were prepared: Pd with an average size of 40 nm (Kyoto University and Kyushu University), BT-Y2c with an average particle size of 200 nm and a cube-like shape (Yamanashi University), and BT-S3 and BT-S5 with average sizes of 300 and 500 nm (Sakai Chemical Industry Co., Ltd.), respectively.

Figure 2(a) shows a reconstructed 3D image of fine BaTiO₃ (BT-S3) particles with a size of ~300 nm. Cross-sectional drawings of the norm and phase, which correspond to electron density and strain, respectively, are also shown. Stress concentration, as indicated by the red circles, is observed at the outer shell of the particle. We could observe the 3D structures of an individual whole single particle, including particle size, shape, and strain within 10-nanometer resolution. Figure 2(b) shows the phase (strain) distribution of a cross-section for a series of samples of various sizes obtained by the Bragg-CDI method. Because of the introduction of the vacuum chamber, as mentioned above, we could measure particles with a wide variety of sizes, from 40 nm to 500 nm. The inhomogeneities of the phase shown in Fig. 2(b) correspond to the strains or phase defects associated with the grain boundaries inside the particles. To clarify this supposition, further investigations on changing the sample environments are required and are now actively under construction.

In summary, the shape and size of the particles were successfully reconstructed. The details of the internal structure and reverse surface of the particles were also obtained. This information is not available from SEM measurements. Our technique can be used to study particles as small as ~ 40 nm in size. Bragg-CDI is an effective technique for investigating nanosized crystalline particles and will open the door to studying particles located within devices, such as MLCCs, which are inaccessible by electron beam techniques.

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Automated stopping of spectral measurements with active learning

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An improvement in experimental efficiency is essential in materials science. The research and development of materials with enhanced functionality and the discovery of novel materials can be accelerated through high-throughput experiments. 'Measurement informatics' is a methodology to enhance experimental efficiency using machine learning techniques. Active learning is a machine learning scheme to obtain predictive models with high precision at a limited cost through the sophisticated selection of samples for labelling. We have previously developed a method to improve the efficiency of Xray magnetic circular dichroism (XMCD) spectral measurements with active learning [1]. XMCD spectroscopy is a derivative experimental technique of X-ray absorption spectroscopy (XAS) that uses circularly polarized X-rays to investigate the elementspecific magnetic properties of materials. The XMCD spectral measurement with active learning depends on the magnetic moment evaluated using the predicted spectrum as the stopping criterion. This condition limits the method to a spectral measurement in which the relationship between the spectrum and physical parameter is known. Moreover, the physical parameters must be evaluated immediately from the spectrum to be included in the active learning cycle of the measurement and Gaussian process regression (GPR). Generally, the evaluation of physical parameters from the spectra is not straightforward; for example, a comparison between the experimental and theoretical spectra is required. To apply spectral measurements with active learning to general types of spectra, a stopping criterion without a physical parameter is required.

In this study, we applied a universal stopping criterion for XAS spectral measurements using active learning [2]. The stopping criterion is based on the stability of expected generalization errors [3]. This stopping criterion is completely evaluated on a mathematical basis; therefore, it is applicable to general spectral measurements, whose relationship with the physical parameter is unknown.

Figure 1 shows a flowchart of a spectral measurement with active learning. Spectral measurement with active learning consists of the following steps: (1) First, initial spectral data $Y_0(X_0)$ are sampled. Here, $Y_0 = (y_i, ..., y_j)$ represents the spectral intensity at energy point $X_0 = (x_i, ..., x_j)$. (2) Subsequently, GPR is applied to the initial data and mean μ_n and standard deviation



Figure 1. Flowchart of the spectral measurement with active learning. T. Ueno *et al.*, npj Comput. Mater. **7**, 139, (2021); licensed under CC BY 4.0. Modified from the original.

 σ_n for each energy point are evaluated. n = 0, 1, ... represents the number of samplings (n = 0 for initial sampling). Mean μ_n is regarded as the predicted spectrum. The stopping criterion is evaluated based on the results of the GPR fitting. (3) If the stopping criterion is not satisfied, then next sampling point x_{next} is automatically determined based on the acquisition function. (4) Subsequently, a new energy point is sampled and added to the measured data, $\mathbf{Y}_n = (y_i, ..., y_{next}, ..., y_j)$. Finally, by repeating process (2), GPR is applied to $\mathbf{Y}_n(\mathbf{X}_n)$. Processes (2)–(4) are repeated until the stopping criterion is satisfied.

Approximating the spectrum by GPR is considered to be a problem in supervised learning. In this setting, the goodness of approximation is evaluated using the average prediction error of the intensity at an unseen energy point, which is called the generalization error. The error ratio is defined using the generalization error as the stopping criterion. Figure 2 shows a schematic of the stopping criterion with the error ratio. Error ratio λ_n is evaluated for each active learning cycle, and the measurement is stopped when λ_n becomes smaller than predetermined threshold λ .

We applied the aforementioned method to simulated noisefree Ni $L_{2,3}$ XAS of Ni²⁺ to verify the effectiveness of the overall strategy. XAS spectra were simulated using CTM4XAS [4]. Figures 3(a)-(f) show snapshots of the GPR fitting of XAS. In Fig. 3(a), randomly selected initial data points and GPR fitting are shown. The covariance function exhibited a large standard deviation between the sampled data points, and the next sampling point was chosen from the x values with a large value of the acquisition function. GPR fitting after several samplings is shown in Fig. 3(b). A rough spectral shape appeared; however, the standard deviation was still large. Figure 3(c) shows the GPR fitting after 50 samplings. Satellite peaks appeared around the L_3 main peak, and multiplet structures appeared around the L_2 peak. Figures 3(d)–(f) show the results for different stopping timings, that is, different thresholds λ . The standard deviation was relatively small compared to the intensities of the L_3 and L_2 peaks in the entire energy range. The intensity relationships of the multiplet structures around the L_2 peak were correctly approximated. In Figs. 3(d) and 3(e), the difference between the GPR fitting for different thresholds appears only in the standard deviations around the non-peak regions. As shown in Fig. 3(f),



Figure 2. Schematic of the stopping criterion with the error ratio. T. Ueno *et al.*, npj Comput. Mater. 7, 139, (2021); licensed under CC BY 4.0.



Figure 3. Gaussian process regression for data size of (a) 10 (initial sampling), (b) 20, (c) 60, (d) 94 (λ =0.1), (e) 116 (λ =0.05), and (f) 178 (λ =0.025). (g) Test error versus data size. (h) Error ratio versus data size. T. Ueno *et al.*, npj Comput. Mater. 7, 139, (2021); licensed under CC BY 4.0. Modified from the original.

the GPR fitting for a data size of 178 was almost the same as the GPR fitting for a data size of 116, but an increase in sampling points around the peak region was observed.

To visualize the progress of the spectral measurement and the stopping timing, the test error and error ratio versus data size are shown in Figs. 3(g) and 3(h), respectively. As shown in Fig. 3(g), the test error steeply decreased with increasing data size as compared to the error ratio in the initial stage of the measurement and converged to a constant value. As shown in Fig. 3(h), the error ratio decreased with increasing data size and converged to an almost constant value. The spikes in the error ratio resulted from the subtraction of the test errors at the steps used to evaluate the upper bound of the test error. The spikes in the error ratio coincided with the spikes in the difference in the test error between the n-th and (n-1)-th sampling. Spikes are inevitable because discontinuous improvement in test errors occasionally arises in active learning. The minimum value of the error ratio is plotted in Fig. 3(h). It was revealed that the minimum value of the error ratio gradually decreased with an increase in the data size. Therefore, immediate early termination of the measurement could be avoided. The vertical dashed lines in Figs. 3(g) and 3(h) indicate the stopping timings for different thresholds corresponding to the GPR fitting shown in Figs. 3(d)-(f). These results indicated that the automated stopping of the XAS measurement based on the generalization error provided GPR fitting with small errors from the ground truth spectrum with a reduced data size.

The XAS spectral measurement with active learning was applied to other spectra, such as simulated Co^{2+} and $Mn^{2+} L_{2,3}$ XAS and experimental Co, Ni, and Mn $L_{2,3}$ XAS for Co and Ni metals and MnO₂ powder. It was concluded that the spectral measurement with active learning was effective for XAS of various elements and with experimental noise.

In summary, we applied a stopping criterion based on the stability of the expected generalization errors for XAS spectral measurements with active learning. This stopping criterion could be evaluated from the self-contained information of GPR fitting. The automated stopping of the XAS spectral measurement was realized with an approximated XAS spectrum with sufficient accuracy. The proposed method can be applied to not only spectral measurements but also other types of measurements. It is expected to enhance experimental efficiency and pave the way for autonomous experimental systems in materials science.

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Theoretical study on the time-resolved structure factor in a photoexcited Mott insulator on a square lattice

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Ultrafast optical pulses are a common tool for investigating novel nonequilibrium phenomena in strongly correlated electron systems. One typical method using this tool is the pump-probe optical technique, which has been widely applied to various strongly correlated materials. The pump-probe technique can detect two-particle excitation but cannot observe momentumdependent collective excitation owing to the negligibly small momentum transfer of probe photons. Conversely, time- and angle-resolved photoemission techniques can probe momentumdependent electronic excitation after pumping; however, they are limited to a single-particle process, where collective excitation appears indirectly.

The recent development of time-resolved resonant inelastic Xray scattering (trRIXS) has opened a new avenue for probing collective two-particle excitation, from which novel photoinduced non-equilibrium phenomena can be investigated in a wide range of momentum and energy spaces. RIXS tuning of incident X-rays for the L edge in transition metals can probe not only charge excitation but also magnetic excitation. Motivated by this development, we theoretically investigated momentumdependent spin dynamics that evolve after pumping within a femtosecond timescale in an antiferromagnetic Mott insulator on a square lattice [1-3]. Characteristic temporal oscillations were demonstrated for the static spin structure factor, exhibiting an antiphase behavior in two orthogonal directions in momentum space: parallel and perpendicular to the electric field of the pump pulse. By analyzing a time-dependent wave function after pumping, we found that the oscillation period was determined by bimagnon excitation in the Mott insulator. This theoretical prediction will be confirmed for Mott insulating cuprates and iridates once trRIXS progresses to a femtosecond timescale.

To describe the Mott insulating states in cuprates, we used a single-band Hubbard Hamiltonian H₀ on a square lattice at half-filling, including the nearest-neighbor (t_h), next-nearest neighbor (t'_h) hopping terms, and on-site Coulomb interaction (U) term. We set U/t_h = 10 and t'_h/t_h = -0.25, which are typical values for cuprates, with t_h~0.35 eV. Hereafter, we use t_h = 1 as the energy unit and 1/t_h as the time unit. An external spatially homogeneous electric field applied along the x-direction via Peierls phase substitution was incorporated in the hopping terms, leading to the time-dependent Hamiltonian H(t). The vector potential A(t) = $A_0 \exp[-(t-t_0)/(2t_d^2)]\cos[w_p(t-t_0)]$ was introduced in the phase, where a Gaussian-like envelope centered at t₀ has a temporal width t_d and a central frequency w_p. We set $A_0 = 0.5$, $t_0 = 0$, $t_d = 0.5$, and $\omega_p = U$.

The time-dependent wave function $|\psi(t)\rangle$ is described by the time-dependent Schrödinger equation. When a time step Δt between two wave functions is as small as $\Delta t = 0.01$, the solution to the Schrödinger equation is given by $|\psi(t+\Delta t)\rangle = \exp[-iH(t)\Delta t]|\psi(t)\rangle$. To calculate the solution, we employed sequential operations of H(t) based on the Taylor expansion: $|\psi(t+\Delta t)\rangle = S_{1=0}|\phi_1\rangle$, where $|\phi_0\rangle = |\psi(t)\rangle$ and $|\phi_{1+1}\rangle = -iH(t)\Delta t$

/(l+1) $|\phi_{l+1}\rangle$. The summation can be truncated if the norm $\langle \phi_M | \phi_M \rangle$ is sufficiently small.

To calculate the dynamic charge and spin correlation functions and the K-edge RIXS spectrum, a real-time representation of the time-dependent cross section has been previously used [4,5]. In this study, we used the spectral representation of dynamic correlation functions, regarding $|\Psi(t)\rangle$ as an initial state, which is easily formulated for systems without time-dependent terms in their Hamiltonians. Applying this formalism to a time region after turning off the pump pulse, t >toff, where the Hamiltonian is time independent, we obtained the time-resolved charge and spin dynamic correlation functions with momentum **q** and frequency ω , denoted by N(**q**, ω ;t) and S(**q**, ω ;t), respectively [1].

Fig. 1 shows the dynamic charge structure factor at t = 5 and N($\mathbf{q}, \omega; t=5$). The factor N(\mathbf{q}, ω) before pumping exhibited momentum-dependent excitations across the Mott gap, which has been observed by Cu K-edge RIXS in insulating cuprates. After pumping, an in-gap excitation emerged inside the Mott gap below $\omega \sim 7$, with a broad spectral weight. The momentum dependence of the in-gap spectral weight was qualitatively similar to that of N(\mathbf{q}, ω) for the two-hole-doped 4 × 4 Hubbard model; however, its center of gravity was clearly higher in energy and its spectral distribution was sharper. These differences demonstrate the contrasting behaviors of chemical doping and photodoping and



Fig. 1 Dynamic charge structure factor before and after pumping for the half-filled 4×4 Hubbard lattice with $t_h = 1$, $t'_h = -0.25$, and U = 10. The black dashed line on each panel represents the equilibrium $N(\mathbf{q}, \omega)$ for \mathbf{q} denoted in the panel. The black and green solid lines represent $N(\mathbf{q}, \omega; t = 5)$ for \mathbf{q} denoted by the same color in each panel.



Fig. 2 Dynamic spin structure factor at t = 5 (a) and t = 7 (b) after pumping for the half-filled 4×4 periodic Hubbard lattice with $t_h = 1$, $t'_h = -0.25$, and U = 10. Momenta **q** are shown in each panel and their color is the same as that of the solid lines.

may be caused by a drastic change in the electronic state owing to photoirradiation.

Fig. 2 shows the dynamic spin structure factor $S(\mathbf{q},\omega;t)$ at t = 5(a) and t = 7 (b). At t = 5, the weight at $(\pi/2,0)$ was slightly larger than that at $(0,\pi/2)$, as shown in Fig. 2(a). Interestingly, the intensity was strongly time-dependent; at t = 7, the intensity at $(\pi/2,0)$ was smaller than that at $(0,\pi/2)$, as shown in Fig. 2(b). This indicates the presence of an oscillation in the spectral intensity as a function of t.

Because the oscillation of the spectral weight causes a change in the integrated intensity with respect to ω , it is useful to investigate the time-resolved static spin-structure factor S(q;t), as shown in Fig. 3. To calculate S(q;t), we introduced various boundary conditions and averaged them to minimize finite-size effects. As presented in Fig. 3, applying a pumping pulse enhanced the magnitude of S(q;t) for small q, whereas this reduced its magnitude for $\mathbf{q} = (\pi, \pi/2)$ and $(\pi/2, \pi)$ as well as for (π,π) (~40 % reduction, but not shown here). This is because the resonant photoirradiation on the antiferromagnetic Mott insulator destroys the short-range antiferromagnetic correlation, leading to the suppression of spin correlation for large q values but enhancement for small \mathbf{q} values. $S(\mathbf{q};t)$ exhibited oscillations, except for $\mathbf{q} = (\pi/2, \pi/2)$; the oscillations were antiphase between two momenta equivalents such as $(\pi/2,0)$ and $(0,\pi/2)$. We found that the antiphase behavior originates from the B1g channel of the two-magnon excitation in the antiferromagnetic Mott insulator [1-3]. We expect that trRIXS will be able to detect such oscillations in the near future.



Fig. 3 Time-resolved static spin structure factor S(q;t) for the half-filled 4×4 Hubbard lattice with $t_h = 1$ and U = 10. The color of the solid lines corresponds to the color of the **q** value denoted in the panel.

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Research Activities of the Institute of Quantum life Science at KPSI

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Institute of Quantum Life Sciences

Quantum life science has emerged to explore new frontiers in biology from the perspective of the quantum domain to describe the dynamics and functions of cells and organisms. The Institute of Quantum Life Science at the National Institutes for Quantum Science and Technology (QST) was established in April 2019 to pioneer a new field in life sciences. Investigations at the Institute of Quantum Life Science are expected to follow two approaches. The first approach is to figure out whether quantum mechanics play an essential role in biological systems. The second approach is to apply quantum science and its technologies for measuring physical and chemical parameters (i.e. temperature, pressure, and pH) inside the cell at an unprecedented minute level, in order to gain information on the structure and dynamics of biomolecules. Using the above two approaches, the goal of the Institute of Quantum Life Science is to discover the essential principles of life and to apply the knowledge developed during the course of investigation in medical sciences and social activities (Fig. 1). The field of quantum life science is envisaged to produce unanticipated discoveries that will expose the existence of quantum phenomena in nature, ultimately leading to their exploitation, which, in turn, would benefit society. At the Kansai Photon Science Institute (KPSI), three research groups in the Institute of Quantum Life Science are carrying out their research by conducting experiments and computer simulations (Fig. 1).

The Molecular Modeling and Simulation Group aims to understand the in vivo function of biomolecules at the atomic level. The group uses computer simulations and bioinformatics based on experimental results, such as crystallographic and scattering data obtained by various types of radiation (x-rays, neutrons, and electrons) and cryo-electron microscopy. The main research target is to determine how the dynamics of protein/DNA/RNA complexes relate to essential cellular functions such as transcription, translation, replication, and repair. One of the focuses of this group is to quantitatively predict the stability of supramolecular complexes such as chromosomes. This year, the research group carried out a simulation to elucidate the overall structure of a protein complex involved in generating circadian rhythms. It is well known that life on Earth has developed circadian rhythms to adapt to daily environmental changes. Cyanobacteria are the simplest organisms with a robust circadian clock; therefore, they have become one of the most useful model organisms for circadian biology. The central oscillator that generates the circadian rhythm in the cyanobacterium Synechococcus elongatus PCC 7942 is comprised of three proteins: KaiA, KaiB, and KaiC. From the experimental results of small-angle x-ray scattering (SAXS) and small-angle neutron scattering (SANS) of the protein complex, the group built an atomic model of the overall structure [1]. The implications of this study will lead to a better understanding of how the cyanobacterial circadian clock works, and thus, will provide fundamental insights into circadian biology. The results have been described in an associated report. The group has also been working on the structural mechanism of NSD2, which

regulates the methylation level of nucleosomal histone H3. Several lines of evidence demonstrate the critical importance of the strict regulation of cellular histone H3 methylation. One significant effect of cellular H3 methylation is that it regulates gene expression by controlling the extent of repressive chromatin domains in genomic DNA. NSD2 is a member of the NSD family that catalyzes the mono- and di-methylation of histone H3, and the deregulation of NSD2 activity is implicated in carcinogenesis. In this study, the structure and dynamics of NSD2 binding to histone H3 were simulated, based on the structure obtained using cryo-electron microscopy [2]. The results indicated that mutations associated with cancer increase the tendency of the autoinhibitory loop to adopt an open state, which provides insight into the regulatory mechanisms of NSD2 proteins under normal and oncogenic states. The details are described in the following report.

The main goal of the DNA Damage Chemistry Group is to clarify the nature of DNA damage induced by various agents, particularly damage from ionizing radiation. The group aims to utilize new experimental techniques that can reveal the structure of DNA damage as well as the recognition of DNA damage by repair enzymes at the nanometer scale. The focus of this group is currently on "clustered DNA damage" in which two or more DNA lesions are located within one to two helical turns of DNA (within several nanometers along the DNA). Clustered DNA damage is considered challenging to repair, and thus is a detrimental type of damage induced by ionizing radiation. However, its presence and microstructure remain elusive, as few experimental methods have been able to obtain data on the spatial distribution of DNA lesions. This research group established a novel approach for measuring the localization level of DNA damage by directly visualizing it. Atomic force microscopy (AFM) has a resolution at the nanometer/sub-nanometer scale, and thus, DNA can be directly visualized. The research group labeled DNA damage (abasic sites) by attaching aldehyde reactive probes with biotin and then attaching streptavidin to biotin. The large molecular size enabled the detection of streptavidin using AFM. Abasic sites were directly induced or further revealed by removing the damaged bases by DNA glycosylases after irradiation. Using this method, the complex nature of clustered DNA damage was visualized for the first time. The research group further developed a promising method for detecting clustered DNA damage using fluorescence resonance energy transfer (FRET). In this method, aldehyde/ketone moieties, such as those at abasic sites in irradiated DNA, are labeled by aminooxyl fluorophores. Fluorescence anisotropy enabled the estimation of the apparent base-pair separations between lesions in a cluster produced by an ion track and revealed that the yield of clustered abasic sites increased with increasing linear energy transfer (LET) of the radiation. Both the AFM and FRET approaches confirmed that these novel analyses have the potential to discover the qualitative and quantitative differences in clustered DNA damage produced by various types of ionizing radiation.

Using cutting-edge quantum technologies and advanced computer simulations, the Institute aims to contribute (1) to the elucidation of the basic principles of life, and in longer term, (2) to the benefit for the everyday life of the society. Groups at KPSI Fast processes(~fsec order),Minute regions(~nm order) Mechanism of **DNA** Damage Molecular Modeling and Chemistry **Mutagenesis** Group Simulation group **Research Group** DNA damage Novel mutation Structure and detection and induction dynamics of damage recognition pathways biomolecules Basic principles of life Medical applications, Benefit for the society

Institute of Quantum Life Science

Figure 1.

The goal of the Mechanism of Mutagenesis Group is to elucidate the underlying mechanisms causing mutations, which are highly relevant to carcinogenesis and the evolution of life. One aspect of this group's research focuses on events at very early stages (around a femtosecond to picosecond) within a space on the scale of nanometers after energy transfer from ionizing radiation, especially from ions. Using Monte Carlo simulations, the research group demonstrated that when water is exposed to densely ionizing ion particles, some of the secondary electrons ejected from water molecules are trapped within the electric potential created by the ionized water molecules. This result led to the realization that the radial dose near the track of a densely ionizing ion particle is much higher than previously understood. This highly localized energy deposition is considered to produce a high yield of clustered DNA damage and thus has important implications for the drastic effect of ion particles on cells. The research group succeeded in arriving at a deeper understanding of the state immediately after energy was deposited near the radiation track by examining the Debye length of the produced plasma [3]. A detailed description is provided in the associated report. Another research interest of this group is the biological effects of cluster ions. They irradiated bacterium (B. subtilis) spores with clustered protons and found that (1) 2 MeV H₂ (clustered proton ions consisting of two) showed a higher lethality per particle than 1 MeV H and (2) 2 MeV H₂ showed a much lower lethality than 340 keV H, which has a similar linear energy transfer (LET) with 2 MeV H₂. The 2 MeV H₂ likely dissociates into a pair of 1 MeV H by losing the bonding electrons at the target surface. This consideration suggests that the deposited energy over the dissociated two protons separated by just several nanometers greatly diminishes the lethal effect. The group also carried out whole-genome sequencing of the irradiated genomes to determine the molecular nature of mutations but failed to find a clear difference between non-clustered and clustered protons. These results suggest that cluster ions are a useful tool for understanding the biological effects resulting from the spatial energy deposited by high LET radiation [4].

Quantum life science is related to the interactions between dynamical phenomena at extremely short time scales and minute length scales, that is, from atto to femtosecond energy transfer processes at the (sub)nanometer scale. Over the long term, the field of quantum life science is expected to produce unparalleled discoveries based on the significance of quantum phenomena in biological systems. To achieve such goals, the research of the three groups of the Institute of Quantum Life Sciences at the KPSI is ongoing.

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Integrative structure modeling of KaiABC clock protein complex

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Every organism on Earth possesses a circadian rhythm that repeats approximately every 24 h and is driven by a circadian clock. The molecular machinery of the circadian clock in cyanobacteria (Kai-clock) comprises three proteins: KaiA, KaiB, and KaiC. These proteins undergo an association–dissociation cycle, coupled with the phosphorylation–dephosphorylation oscillation of KaiC in the presence of adenosine triphosphate (ATP)^{1,2}. During the circadian cycle, the Kai-clock system generates three forms of complexes, two binary complexes KaiAC and KaiBC, and one ternary complex KaiABC, at specific clock phases^{3,4} (Fig. 1).

KaiA comprises an N-terminal domain (NA), a canonical linker, and a C-terminal domain (cA), and forms a homodimer (A₂) through cA^{5,6}. KaiB assumes a single thioredoxin domain and forms a homotetramer (B4)^{7,8}. KaiC comprises two domains (CI and CII) and forms a homohexamer (C_6) with a double doughnut-like shape9-11. The binary complex KaiAC (A2C6) is formed by an A2 dimer and a KaiC hexamer through interactions at the C-terminal tails of the CII domains¹²⁻¹⁴. Another binary complex, KaiBC (B₆C₆), is formed by a KaiC hexamer and six KaiB molecules arranged on top of CI domains^{3,15,16}. Regarding the ternary complex KaiABC, the supramolecular architecture and stoichiometry of the complex have been controversial for a long time¹⁷⁻¹⁹. Recently, a cryo-EM (cryo-electron microscopy) study revealed the structure of the complex (A12B6C6), in which six A₂ dimers were captured onto the KaiB ring in the B₆C₆ subcomplex^{20,21}. However, the NA domains of the A₂ dimers were absent from the structure, suggesting that the domains could fluctuate in the complex. Previously reported mutational studies have indicated that the NA domain is essential for the generation of circadian rhythms^{22,23}. Therefore, it is crucial to determine the overall structure of the A12B6C6 complex, including the locations of NA domains.

Small-angle X-ray and neutron scattering (SAXS and SANS) provide structural information regarding supramolecular complexes in solution²⁴⁻²⁶. We utilized SAXS coupled with size-exclusion chromatography (SEC–SAXS)²⁷⁻²⁹ to avoid scattering from undesirable components and inverse contrast-matching SANS (iCM-SANS) to selectively observe scattering originating from KaiA in the A₁₂B₆C₆ complex.

We established candidate models of the $A_{12}B_6C_6$ complex computationally, which can reproduce the results of SAXS and SANS (see Fig. 2). We used the crystal structures of a KaiC hexamer (C₆, PDB code: 3dvl)¹¹ and a ternary complex (_CA₂-B-C_IC, PDB code: 5jwr)²¹ comprising two C-terminal domains of the KaiA dimer (_CA₂), a KaiB monomer (B), and a CI domain of KaiC (_{CI}C) to build a model of the cA₁₂B₆C₆ complex (A₁₂B₆C₆ complex without NA domains), which agreed well with the previously reported cryo-EM structure. Subsequently, we combined it with the crystal structure of a full-length KaiA dimer (A₂, PDB code: 1r8j)²⁰ to obtain the overall structure of the A₁₂B₆C₆ complex. However, in this structure, the NA domains overlapped structurally with KaiB, indicating that spatial rearrangements of NA domains were necessary. To remove this



Figure 1: Proposed oscillation scheme for cyanobacterial circadian clock system (Kai clock system). Blue squares, orange triangles, and green spheres represent monomers of KaiA, KaiB, and KaiC, respectively. Two phosphorylated sites, i.e., Ser431 and Thr432, exist in CII domain of KaiC. Phosphorylation cycle is as follows: $S/T \rightarrow S/pT \rightarrow pS/pT \rightarrow pS/T \rightarrow S/T$ (p indicates phosphorylated residue). Phosphorylation–dephosphorylation cycle of KaiC is coupled with its association and dissociation with KaiA and KaiB.



Figure 2: Modeling procedure of overall structure of $A_{12}B_6C_6$ complex.

overlap, we systematically altered the positions and orientations of the individual NA domains. At this stage, we disregarded the linkers connecting the NA and cA domains to reduce the computational cost. Finally, we constructed ~ 20 million models of the A₁₂B₆C₆ complex as the initial structural candidates.

We calculated the scattering curves for each model and selected models that indicated small χ^2 values for both SAXS and SANS. Interestingly, three different types of models agreed well with the SAXS results, but only one of them agreed with the SANS results (Fig. 2). This shows that the combination of multiple experiments is important for structural determination.

After modeling the linkers connecting the NA and cA domains, which were not considered in the previous steps, we performed MD simulations on the models that agreed well with both the SAXS and SANS results to verify their stability in water. This process was necessary because the protein domains were regarded as rigid bodies, and only the exclusive volume of the molecules was considered. In fact, most of the models (> 90%) were unstable and could not maintain small χ^2 values for SAXS and SANS during the simulations; this indicates that the MD simulation may be an effective tool for model selection.

In summary, this study demonstrated that the integrated approach of modern solution scattering methods, i.e., SEC– SAXS and iCM-SANS, as well as computational modeling and molecular dynamics simulation can effectively resolve the structures of supramolecular complexes harboring dynamically fluctuating domains/subunits, such as the KaiABC complex.

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Oncogenic mutations E1099K and T1150A enhance NSD2 methyltransferase activity by increasing the auto-inhibitory loop dynamics

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Histone post-translational modifications (PTMs) are distributed throughout chromatin in a context-dependent manner. It is known to regulate diverse genome-related and downstream processes. Alterations, either in the dynamics or distribution of histone PTMs, alter the gene expression profiles of cells. These disturbances are often linked to the development of various pathological conditions in humans [1]. Dimethylation of histone 3 lysine 36 (H3K36me2) is one such PTM, catalyzed by the NSD family member protein NSD2 (also known as WHSC1/MMSET). It incorporates mono- and dimethylation at lysine 36 of histone 3 (H3K36). Aberrant upregulation of H3K36me2 by induced NSD2 overexpression or due to NSD2 point mutations (E1099K and T1150A) is associated with multiple myeloma, lymphoblastic leukemia, and other types of cancers and developmental anomalies in neurons and cardiovascular tissues.



Figure 1. NSD2-nucleosome complex. **a** NSD2 bound to nucleosome with partially unwrapped DNA near the entry/exit site. **b** H3-bound and H3-free NSD2 structure comparison. **c** Comparison of autoinhibitory loop conformation in H3-free (blue) and H3-bound NSD2 (orange). **d** Hydrophobic residues surrounding the H3V35-binding hydrophobic cavity showing the overlap between the positioning of L1184 in H3-free and V35 in H3-bound NSD2 structure.

Biochemical studies have shown that NSD2 has a strong specificity for nucleosomal substrates. Our nucleosome-bound NSD2 cryo-EM structure revealed that nucleosome-NSD2 association requires the removal of one DNA gyre near the entry/exit site to access the H3K36 located between the two DNA gyres in the canonical nucleosome. This leads to the partial unwrapping of DNA at the nucleosomal entry/exit. In this binding mode, NSD2 establishes extensive interactions with the H3 N-terminal tail, H2A C-terminal portion, H3 first α -helix, nucleosomal DNA, and external linker DNA (**Fig. 1a**) [2].



Figure 2. Cross-correlation plots of the residue-wise fluctuation. See the legend in Figure 3 for the definitions of R1, R2 and R3.

The overall structures of the H3-bound and H3-free NSD2 were similar (**Fig. 1b**). However, compared to H3-free NSD2, where the autoinhibitory loop in the SET domain occupies the substrate-binding cleft, assuming a closed conformation, leaving no room for H3 tail accommodation, in the H3-bound NSD2 structure, the autoinhibitory loop adopts an open conformation. (**Fig. 1c**). Point mutations E1099K and T1150A increase NSD2 catalytic activity without increasing its affinity for nucleosomes [2]. There remains a gap in the understanding of the NSD2 catalytic enhancement mechanism following these two point mutations.

To fill this gap, we carried out MD simulations of the H3 free NSD2 wild type and mutants. The NSD2 structure was extracted from the nucleosome-bound complex and used as the starting point for the simulation [2]. Comparative analysis of the MD trajectories based on time-dependent changes in secondary structures and residue-wise contact maps clearly showed that NSD2 wild type, E1099K, T1150A, and the double mutant E1099K-T1150A structures were stable and maintains similar

global conformations, except in the autoinhibitory loop. The autoinhibitory loop remains flexible in all systems [2]. The dynamic cross-correlation analysis of the wild type NSD2 SET domain residue-wise fluctuations showed anti-correlated motions among residues in the regions R1-R2 and R1-R3 (Fig. 2), which are absent in the mutants. The absence of anti-correlated motion suggested that the movement of R2 and R3 was independent of R1 in the NSD2 mutants (Fig. 2). R1 contained the mutation site E1099 and residues forming the H3V35- and H3P38-binding hydrophobic patches, whereas R2 and R3 constituted the mutation site T1150A and the autoinhibitory loop, respectively (Fig. 1d and 3a).

Independent movement of R3 suggests the possibility of autoinhibitory loop adopting an open conformation more frequently, facilitating H3 tail binding to NSD2 mutants. For validation, the open-close motion of the autoinhibitory loop was quantified based on four distance-based locks as shown in **Fig. 3b**. It was observed that the autoinhibitory loop tends to assume an open conformation far more frequently in mutants (9.7%, 6.3%, and 18.8% in the case of E1099K, T1150A, and E1099K-T1150A, respectively) than in the wild type (only 0.86%) (**Fig. 3c**).



Figure 3. Open-close dynamics of the autoinhibitory loop. a Regions R1, R2, and R3 are shown on the NSD2 SET domain structure. b Def. of the four distance-based locks to quantify the autoinhibitory open-close dynamics. c Table showing the percentage of open autoinhibitory loop conformation in NSD2 wild type and mutant.

A detailed examination of MD trajectories revealed the importance of the interaction between the residue L1181 and the hydrophobic patches (especially T1121 and T1150), whose replacement by alanine leads to NSD2 hyperactivity. Based on the MD simulation results, a working model of NSD2 aberrant catalytic enhancement was proposed. In the wild type H3-free NSD2, the autoinhibitory loop exhibits multiple conformations. However, the hydrophobic interactions between residues L1181, C1183 and L1184 and the hydrophobic patches maintained the autoinhibitory loop in a closed conformation, leaving no space for H3 tail binding. Mutations E1099K and T1150A affect the conformational dynamics of the autoinhibitory loop and promote its tendency to adopt an open conformation, leaving sufficient room for H3 tail-binding. A possible mechanism by which point mutations affect the autoinhibitory loop conformation could be via the network of salt bridge interactions at the base of the H3

binding pocket surrounding the loop (**Fig. 4**). Mutations E1099K and E1099K-T1150A seem to remodel the salt bridge interaction network in the NSD2 wild type by adding a novel salt bridge with D1125, accompanied by a decrease in salt bridge formation between D1123 and K1152. Similarly, in the T1150A mutant, the D1123-K1152 salt bridge formation was seen to decrease (**Fig. 4**).



Figure 4. Schematic representation of the residues forming the salt bridge network around the autoinhibitory loop. Dashed lines depict salt bridges between residue pairs with the percentile of bridge formation.

The D1123-K1152 salt bridge connects two β -sheets that form the H3V35 and H3P38 binding patch. Interestingly, the autoinhibitory loop opens when the D1123-K1152 salt bridge is lost. In summary, our study provides insight into the mechanism of catalytic enhancement of NSD2 following oncogenic mutations E1099K and T1150A and facilitates the development of inhibitors against these proteins for cancer treatment.

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A Debye shield formed by track potential and transport of secondary electrons under heavy ion irradiation

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1. Introduction

Under heavy ion irradiation, a special physical phenomenon appears near the incident ion path (IP); that is, a strong electric field created near the IP affects the transport of secondary electrons (SEs). A SE is an electron created by incident impact ionization, and the potential created by this electric field is called "track potential (TP)". This effect is expected to form plasma [1], which contributes to increasing the local dose in this region. The plasma formed by heavy-ion irradiation is assumed to be important; however, this plasma has seldom been studied. To the best of our knowledge, this is because there were no methods to study the plasma produced near the IP before our simulation model was developed [1,2]. Only our simulations [1] succeeded in reproducing the measurement trend of SE yields (Y_{se}) [3]. In this study, we analyzed when, where, and how plasma is created through simulations and Debye shield theory. Furthermore, we studied the effect of this shield on the transport of SEs and TP. The Debye shield decreased the electric field outside the shield itself.

In heavy-ion beam science, SE transport is crucial because SEs transport most of the energy deposited from the incident ion over a wide area. To simulate this transport phenomenon, two types of models were proposed in the early 20th century: freeelectron gas and binary collision. In the free-electron gas model, electron plasma, which is estimated to enhance the local dose near the IP, is created. However, (i) the Debye shield is not considered in this model, (ii) it is difficult to incorporate it in a simulation model, and (iii) free-electron gas is far from the real target. In contrast, the binary collision model is easy to employ in large-scale simulations; however, a plasma is not created in this model.

In our simulation model developed based on a binary collision model, we incorporated the effect of the TP on SE transport [1]. We detected from previous simulations [1] that this effect is expected to produce plasma, which increases the local dose near the IP. A similar increase also appeared in free-electron gas models, as mentioned above. We compared our simulations regarding the probability (P_{set}) of SEs escaping TP [1] with the measurement data of the secondary electron yield (Y_{se}) from the insulator KCl target [3]. Note that this trend cannot be explained by the two aforementioned conventional models because TP is ignored in the binary collision model and production rate of electron plasma in SEs is established regardless of the ion impact ionization cross-section σ in the free-electron gas models. Furthermore, we study the Debye shield theory and compare the simulation results with the analytical results.

2. Simulation model

In conventional simulation models based on the binary collision theory, a continuum is employed as the target. To consider a more realistic transport of SEs, we treated individual water molecules in a rectangular target. The procedures followed by the proposed model are summarized below. Using a random number (N_{ran}), the positions of water molecules are randomly determined to obtain N_{mol} in water. Furthermore, we prepared several targets with different molecular locations. For each target, we simulated a heavy ion passing through. The average values of these results are presented in section 4.

A heavy ion is assumed to enter the target perpendicular to its bottom area. Water molecular ions and SEs are produced according to σ , the center of which is located at the center of each molecule. The initial energy and emitted angle of SEs are determined by fitting the measurement functions. To incorporate the effect of TP on the transport of SEs, we considered the Coulomb interaction of the SEs with molecular ions (F_{ei}) produced by the incident ion impact ionization and other SEs (F_{ee}). In the binary collision model, both F_{ei} and F_{ee} are not considered; however, in the free electron gas model, F_{ei} is considered.

3. Debye shield theory

Using one dimensional Poisson's equation, we obtain

$$n_{\rm e}(r) \sim n_{\rm e0} \ 1/2\pi r \exp(-r/\lambda_D), \tag{1}$$

where n_e , n_{e0} , r, and λ_D are the electron density, constant value, the distance from the ion path, and Debye length, respectively. Using $n_e(r)$, as stated in eq. (1), and ion density, the probability (P_{set}) of the secondary electrons escaping the TP is solved using the following relation:

$$P_{set} \vdash \frac{[A + (\sqrt{S} - A)\exp(-\frac{\sqrt{S}r_e}{C})]}{\sqrt{S}} \quad . \tag{2}$$

Here, r_e is the point at which it is assumed that SEs escape the TP.

4. Results and discussion

Figure 1 depicts $N_{\rm SE}(r)$ vs. r at $E_{\rm ion} \sim 500$ keV/u for $\sigma \sim (a) 8 \times 10^{-16}$ and (b) 2×10^{-15} with $N_{\rm SE}(r) \propto 2\pi r n_{\rm e}(r)$, where $E_{\rm ion}$ denotes the incident-ion energy. In figure 1, functions $N_{\rm SE0} \exp(-r/\lambda_{\rm D})$ are also shown, where $N_{\rm SE0}$ and $\lambda_{\rm D}$ are approximately estimated from our simulation results. As shown in figure 1, $N_{\rm SE}(r)$ almost coincides with the solution of the Poisson's equation.

Figure 2 depicts P_{set} vs. σ at $r_e \sim 2$ and 5 nm, and $t \sim 2$, 3, 4, 5, 10, 40, 80 fs at $E_{\text{ion}} \sim 500$ keV/u. In this figure, the function on the right-hand side of eq. (2) is also plotted. Observe from figure 2 that P_{set} almost coincides with eq. (2). Namely, we succeeded in deriving an analytical solution that approximately reproduces the simulation results of P_{set} . At $r_e < 2$ nm, the term exp ($-\sqrt{\sigma}r_e/C$) in Eq.(2) cannot be ignored for all σ in question because r_e and λ_D are comparable. At $r_e \ge 5$ nm, this term becomes almost zero; therefore, $P_{\text{set}} \propto 1/\sqrt{\sigma}$.



Fig.1 $N_{\text{SE}}(r)$ vs. r at $E_{\text{ion}} \sim 500$ keV/u for $\sigma = (a) 8 \times 10^{-16}$, and (b) 2×10^{-15} , and (d) 4×10^{-15} cm². The simulation results are indicated by the symbol \diamondsuit , whereas the dashed lines represent the functions $N_{\text{SE0}} \exp(-r/\lambda_{\text{D}})$.

5. Summary

In this study, we developed a simulation model that forms the Debye shield. Based on the simulation results of P_{set} , we demonstrated when, where, and how the Debye shield in plasma is created by heavy-ion irradiation. The simulated Debye shield is approximately reproduced by the analytical solution of Poisson's equation. We derived a formula that approximately reproduces the simulation results of P_{set} . This formula is also expressed as a function of only one parameter, namely, σ .



Fig. 2 P_{set} vs. σ for $r_e \sim (a)$ 2 and (b) 5 nm at $E_{\text{ion}} \sim 500 \text{ keV/u}$ and $t \sim 2$, 3, 4, 5, 10, 40, and 80 fs (which are indicated by the symbols \bigcirc , +, -, \times , \diamondsuit , \square , and \triangle , respectively). Thick-solid and dashed lines express functions of the right side of eq. (2) and functions showing the trend decreasing (σ)^{-1/2}, respectively.

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5. 分子科学会優秀講演賞.

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押目 典宏 「コヒーレントX線回折を利用したナノ結晶の3次元イメージング」及び論 文 "Bragg coherent diffraction imaging allowing simultaneous retrieval of threedimensional shape and strain distribution for 40–500 nm particles" 強誘電体会議, 2022-06-01 (2021 年度論文に対する表彰)

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内海 伶那 「Fe-Mo 合金の高温高圧水素化反応の組成依存性」 水素若手研究会, 2021-09-07

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一般社団法人 大阪ニュークリアサイエンス協会, 2021-06-10

11. 泰山賞.

桐山 博光、Alexander Selgeivich Pirozhkov、神門 正城 「世界最高高性能ペタワットレーザ ーの開発と量子ビーム制御」

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 中新 信彦,神門 正城,黄 開,近藤 康太郎
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6. 磁化測定方法.

稲見 俊哉, 菅原 健人, 綿貫 徹, 小出 明広, 野村 拓司 PCT/JP2021/044470 (2021-12-03)

7. 磁化測定装置及び磁化測定方法.

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- 8. データ取得装置、データ補正装置、データ補正方法、プログラム及び記録媒体. 佐々木 拓生, 高橋 正光 特願 2022-041838 (2022-03-16)
- 9. 水素吸蔵材およびその製造方法.

齋藤 寛之, 内海 伶那, 綿貫 徹 特願 2022- 39521 (2022-03-14)

10. 新規ヒトインターロイキンー18変異体及びその用途. 田中 義正, 櫻庭 俊 特願 2021-01941 (2021-02-10)
The Kids' Science Museum of Photons

きっづ光科学館ふおとんは、2020年度の300人を上回る757人の方々にご来館いただきました。

2021 年度も新型コロナウィルスの感染拡大防止のため年度当初より臨時休館することとなりました が、4月に一般社団法人 VR 革新機構(VRIO)の公益事業プログラム(コロナ禍の博物館等を支援する 3D+VR ボランティア撮影)により、科学館内部を撮影していただきました。レーザースキャナーや赤外 線カメラによるデータも用いて構成される 3D ビューコンテンツにより、館内見学を仮想体験できます。 また、超長尺三脚を駆使して計測・撮影されたデータや画像も併用・統合して構築された「ドールハウ ス表示」により、館内部を俯瞰することもできます。

公開 URL https://my.matterport.com/show/?m=wAeZTtF8jv8



科学館 3D ビューコンテンツのドールハウス表示

7 月上旬より課外授業(学習投影)等の場として利用して いただくため、プラネタリウムの上映を事前予約制で再開し ました。7月11日に開館20周年を迎え、7月25日には入館 者数が75万人に達しました。その後、8月半ばに京都府に緊 急事態宣言が発出されたことにともない9月末まで休館、10 月より再開し、プラネタリウムに加え希望者には親子工作を 体験していただいています。偏光板工作や分光板工作のサン プルや作品の試作を行うとともに、ウェルカムボードや月替 わりの偏光グラフィックスの作成・展示も行いました。

関西研の施設公開(10月31日)では、プラネタリウムの 上映を入替制で4回行い、「案内・誘導・上映・座席消毒」の 工程も4セットとなり、慌ただしくも活気のある一日となり ました。



ふおとん 20 周年(7 月 11 日)









偏光グラフィックス展示



75 万人達成(7 月 25 日)

工作体験

2022年1月より、「光の不思議体験コーナー」を受付エリアに開設し、鏡とハーフミラーの間に LED ライトを配置してできる無限鏡(光のトンネル)や錐体万華鏡、立方体万華鏡を体験していただきました。



無限鏡

錐体万華鏡

立方体万華鏡

年度末には、光の科学ゾーンの展示装置「波のテーブル」の改修ならびに館外で記念撮影などに用いるパネルの更新を行いました。



「波のテーブル」



記念撮影パネル

Appendix

共同研究課題、施設共用課題

1) 木津地区

【共同研究課題】

共同研究先	共同研究課題名	担当研究グループ
日本原子力研究開発機構	中性子散乱による生体高分子のダイナミクス解析 に関する研究	生体分子シミュレー ショングループ
大阪大学、東北大学	シンチレータを用いた相対論的イオン検出手法の 開発	先端レーザー技術開発グループ
東京大学	半古典 Vlasov 方程式に基づくレーザー加工シミュ レータの研究開発	超高速光物性研究グ ループ
京都大学	レーザー誘起弾性波技術によるインフラ構造物診 断の高度化に関する研究	X線レーザー研究グ ループ
日本電信電話株式会社	アト秒パルスの波形計測と時間分解計測に関わる 基盤技術開発	超高速光物性研究グ ループ
大阪大学	超高強度レーザーと固体物質からのガンマ線発生 及び核物理に関する研究	高強度レーザー科学 研究グループ
慶應義塾大学、近畿大学	レーザーを活用した整形外科インプラント設置強 度評価機構の開発	X線レーザー研究グ ループ
株式会社島津製作所	高耐性光学デバイスの開発研究	X線レーザー研究グ ループ
宇都宮大学、広島大学、東北大 学	高輝度軟X線発生に関する基礎研究	X線レーザー研究グ ループ
京都大学	TW 級チタンサファイアレーザーを用いた量子ビ ーム発生に関する研究	X線レーザー研究グ ループ
奈良女子大学	レーザー誘起振動波計測の機械学習手法の適用に 関する研究	X線レーザー研究グ ループ
株式会社フォトンラボ	レーザー誘起振動波計測技術の小型化に関する研 究	X線レーザー研究グ ループ
芝浦工業大学	レーザー誘起振動波計測技術の高度化に関する研 究	X線レーザー研究グ ループ
NTT アドバンステクノロジ株 式会社	高耐力軟X線光学素子に関する基礎研究	X線レーザー研究グ ループ
東京学芸大学	リラクサー強誘電体のドメイン形成に関する研究	X線レーザー研究グ ループ
ライトタッチテクノロジー株 式会社	中赤外レーザーを用いた医療応用計測技術の開発	光量子科学研究部

······································	テラヘルツパルス光源による高分子高次構造の制	超高速光物性研究グ
理化子研究別	御	ループ
十四十学	高強度テラヘルツパルス光源による新規物質創成	超高速光物性研究グ
入II风入子	と新規物性発現に関する研究	ループ
大百开册刘兴开发上兴应上兴	有機半導体におけるフォノン・励起子相互作用を利	超高速光物性研究グ
佘 民 无	用した状態制御に関する基盤技術開発	ループ
ら虎眼寺特フ始医病キング	量子メスに向けた粒子線照射計測及び制御技術に	高強度レーザー科学
- 共単県立私士禄医療センター	関する研究	研究グループ
古北上兴 白油制作正	伝典此及始始によってもなってあっている。	先端レーザー技術開
· 果北入子、 局律聚作所	極端系外線頃或の低入射角凹折格士の開発	発グループ
同十九上兴	フェムト秒パルスレーザーの位相受動整合コヒー	先端レーザー技術開
问志仁人子	レント加算に関する研究	発グループ
神戸大学、名古屋大学、東京大	固体飛跡検出器を用いた高エネルギーイオン検出	先端レーザー技術開
学	手法開発	発グループ
· 古如-L 学	高強度レーザーと構造性媒質との相互作用に関す	先端レーザー技術開
京都大学	る研究	発グループ
	宇宙物理の手法に基づく粒子加速・電磁放射に関す	先端レーザー技術開
人阪人子、育山子阮人子	る研究	発グループ
核融合科学研究所、京都工芸	「「「「「「」」」」、「「」」、「」、「」、「」、「」、「」、「」、「」、「」、	X線レーザー研究グ
繊維大学、富山高等専門学校	疑脳相の重士・万士動力子計鼻に関する研先	ループ
十四十岁	1. 此, 承了加速の空空心, 研究, 研究,	高強度レーザー科学
入版入子	レーリー电子加速の女だ化・住能計価研究	研究グループ
住友重機械工業株式会社	量子メス用レーザー加速入射器の評価手法の開発	光量子科学研究部
	量子メス用レーザー加速入射器のターゲット駆動	
日立造船株式会社	装置に関する研究開発	光量子科学研究部
	■ 子戸 日本	
日本電子	開発	光量子科学研究部
	・ 光・電子融合第一原理ソフトウェア「SALMON」の	超高速光物性研究グ
筑波大学	開発及び利用研究	ループ
		高強度レーザー科学
株式会社 NAT	小型レーザー駆動量子ビームの安定運用の研究	研究グループ
	レーザー電子加速に資する防振・光軸安定化に関す	高強度レーザー科学
株式会社日本防振工業	る研究	研究グループ
	フェムト秒レーザー・物質相互作用の高速計測に関	高強度レーザー科学
大阪大字		

【施設共用課題】

課題番号	利用 区分	施設装置	利用課題
2021B-K01	公開	J-KAREN-P レーザー装置	陽子線計測の統計的有意性に基づいたレーザー駆動イ オン加速の高強度化

2) 播磨地区

【共同研究課題】

共同研究先	共同研究課題名	担当研究グループ
物質・材料研究機構	原子二体分布関数法による機能性材料の先進的ナ ノ構造研究	高圧・応力科学研究 グループ
京都大学	同位体特定による局所状態解明のための先進的メ スバウアー分光法の開発研究	磁性科学研究 グループ
京都大学、分子科学研究所、理 化学研究所	コヒーレント X 線を利用したナノ粒子計測に関す る研究	コヒーレント X 線 利用研究グループ
広島大学	コヒーレントX線を利用した強誘電体一粒子計測	コヒーレント X 線 利用研究グループ
京都大学	ナノ粒子を取り込んだがん細胞に及ぼす高エネル ギー単色X線照射の影響に関する研究	高圧・応力科学研究 グループ
JEF テクノリサーチ株式会社	磁気円偏光発光を用いた方向性珪素鋼板の内部磁 区観察のための研究開発	磁性科学研究 グループ
日本原子力研究開発機構	放射光による物質科学に関する研究	放射光科学研究セン ター
産業技術総合研究所	水素雰囲気その場・時分割放射光 X 線全散乱を利 用した水素吸蔵合金の構造変化の観測	高圧・応力科学研究 グループ
東北大学、上智大学	水素雰囲気その場・時分割放射光 X 線全散乱を利 用した水素吸蔵合金の構造変化の観測	磁性科学研究 グループ

【施設共用課題】

播磨地区では 2012 年度より文部科学省のナノテクノロジープラットフォーム事業を受託してお り、放射光科学研究施設を成果公開型課題で利用する外部研究者に対して、特に専用ビームライン における研究支援を強化しています。課題は年 2 回、SPring-8 を運営する公益財団法人高輝度光科 学研究センターの一般課題募集時期(5月及び11月)に合わせて募集しています。

課題番号	利用 区分	利用装置	研究課題
2021A-H01	公開	放射光メスバウアー 分光装置	Dy 放射光メスバウアー吸収分光法の高度化による単分子 磁石の磁気緩和過程解明
2021A-H02	公開	放射光メスバウアー 分光装置	電荷秩序配列に起因する純核ブラッグ散乱の測定
2021A-H03	公開	放射光メスバウアー 分光装置	放射光メスバウアー分光を利用した (A1, Fe3+)00H中の 鉄のスピン状態の圧力依存性の解明
2021A-H04	公開	共鳴X線非弾性散乱 装置	Investigating the electronic structure of Cu2ZnSnSe4 by resonant inelastic x-ray scattering at Cu K-edge
2021A-H05	公開	表面X線回折計	RF-MBE法による窒化物半導体成長初期過程のXRDその場観 察
2021A-H06	公開	表面X線回折計	アモルファスカーボン量の最適化によるグラフェン析出 成長の過飽和度制御
2021A-H07	公開	表面X線回折計	InGaN自立基板製作に向けたその場X線回折測定による成 長制御
2021A-H08	公開	表面X線回折計	グラフェンを用いたサファイア上でのGaNのリモートエピ タキシーのリアルタイムX線回折
2021A-H09	公開	高温高圧プレス装置	ポリエチレンの高温高圧Hexagonal相の存在領域の研究
2021A-H10	公開	高温高圧プレス装置	Colossal Negative Thermal Expansion in Polar Perovskites with Large Tetragonality
2021A-H11	公開	高温高圧プレス装置	放射光透過ラウエによるマグネシウム系材料の曲げ加工 後の内部損傷評価
2021A-H12	公開	高温高圧プレス装置	高圧合成技術を活用した熱化学式水素圧縮機用新規水素 吸蔵合金の探索
2021A-H13	公開	高温高圧プレス装置	高圧下水素化技術による鉄鋼材料の新奇無拡散相の探索 (2)
2021A-H14	公開	高温高圧プレス装置	マグネシウム系遷移金属錯体水素化物における高温高圧 合成過程のその場観察
2021A-H15	公開	高温高圧プレス装置	白色X線と二重露光法による溶接底部の残留応力評価
2021A-H16	公開	高温高圧プレス装置	合金化溶融亜鉛メッキ鋼板抵抗スポット溶接部における 溶接割れの放射光X線イメージング
2021A-H17	公開	ダイヤモンド アンビルセル回折計	PbFe03における圧力誘起金属間電荷移動転移の観測と負 熱膨張化への展開
2021A-H19	非公開	共鳴X線非弾性散乱 装置	RIXSを駆使したSCM415鋼の水素トラップ現象解明

9091 D-U 01	心問	放射光メスバウアー	金属-窒素-ドープカーボン触媒の局所活性サイト構造の	
20210-001	公開	分光装置	解明	
2021B.U02	心問	放射光メスバウアー	真水妻田力下におけるい水妻化物の雪子骨能分析	
2021D 1103	ム田	分光装置	同八米几777 (C37) ③20小米化初99电1 八惠万州	
2021B-H04	公開	放射光メスバウアー	純核ブラッグ散乱法による自然鉄試料のサイト選択的ス	
2021D 1104	公用	分光装置	ペクトル測定	
9091 D-U 05	心問	共鳴非弾性X線散乱	準静的磁場変化における方向性電磁鋼板の内部磁区構造	
20210-000	公用	装置	調査	
2021 B-H OG	心問	丰田V绚同坵卦	放射光X線を用いたヘテロ界面上の転位生成ダイナミクス	
2021D 1100	公用	衣田A林田17日	と転位すべり面分布の測定	
2021 B.H 07	公開	志 ∀ 線同 折 卦	放射光X線を用いた格子不整合III-V族化合物半導体混晶	
20210-007	公用	衣田A林田17日	のリアルタイム構造解析	
9091D-U09	心問	丰田V绚同坵卦	GaN/グラフェン基板上でのInGaNヘテロエピタキシャル成	
20210-000	公用	衣田A林田17日	長における応力緩和過程の観察	
9091D-1100	八月月	表示V绚同长到	InGaNエピタキシャル成長におけるその場X線回折測定に	
20210-009	公開	表面X禄回折計	よる歪み制御	
9091D-1110	心問	吉 润 百 匹 プ レ フ 壮 罢	高圧合成技術を活用した熱化学式水素圧縮機用新規水素	
2021D-H10	ゴ田	同価同圧ノレハ表則	吸蔵合金の水素放出挙動の観察	
2021B-H11	公開	高温高圧プレス装置	水素貯蔵合金の潜在化された水素吸蔵サイトの解明	
9091D-1119	八胆	吉 泪 吉 広 プ レ フ 壮 罟	Mg-W系遷移金属錯体水素化物の高温高圧合成過程のその	
2021D-H12	公開	同価同圧ノレハ表則	場観察	
9091 D-U 19	心問	言泪言にプレッ壮署	中Mn鋼の残留オーステナイト変態挙動,引張変形時の相応	
2021D-H13	公用	同価同江ノレハ表直	力分配挙動と優れた延性発現機構の解明	
9091 D-U 14	心問	吉 润 古 広 プ レ フ 壮 罢	X線回折とX線透過法を用いたTi-Mg系の高圧高温状態のそ	
20210-П14	公開	同価同圧ノレハ表則	の場観察	
2021B-H15	公開	高温高圧プレス装置	高温高圧法を用いた酸水素化物合成の直接観察	
2021B-H16	公開	高温高圧プレス装置	鉄鋼材料の高温圧縮過程のX線その場観察	
0001D 1117	八日日	古泪古口プレッ壮平	放射光白色X線を用いた高強度鋼板抵抗スポット溶接継手	
2021B-H17	公開	向価向圧ノレム装直	の内部ひずみ分布の測定	
0001D 1110	八日日	ダイヤモンド	PbFe03における圧力誘起金属間電荷移動転移の観測と負	
2021B-H19	公開	アンビルセル回折計	熱膨張化への展開II	
2021B-H23	非公盟	大型X線回折計	円筒雪池かしめ部の残留広力評価	
20210 1120	ットムア			

関西光科学研究所での各種シンポジウム・施設公開・出展・アウトリーチ活動

全世界的な新型コロナウイルス感染症 (COVID-19) の影響で、 2021 年度も対面での活動が大きく制限されました。そのような 状況下ですが、関西研(木津地区、播磨地区)の各種シンポジウ ム、セミナー、研究会等について主なものを記載します。

年1回の施設公開について木津地区では2021年10月31日 (日)に人数制限・事前登録のうえ当日の感染症対策を徹底して 開催しました。なお、播磨地区での開催は見送りました。



関西研施設公開(木津地区) スーパーサイエンスセミナー

【木津地区】

1	22-JUN-2021	オンライン開催	主催:阪大・レーザー科学研究所、量研・関 西光科学研究所
	光・量子ビーム科学会	合同シンポジウム 2021	•
	OPTO2020 Symposiu	m on Photon and Beam Science	
2	30-JUN-2021	オンライン開催(発表)	主催:関西文化学術研究都市推進機構
	けいはんな若手研究者	皆交流会(第 44 回)	
3	29-OCT-2021	オンライン開催 (発表)	主催: 関西文化学術研究都市推進機構
	けいはんな若手研究者	皆交流会(第 45 回)	
4	31-OCT-2021	関西光科学研究所(木津地区:京都府木津川市)	主催:関西光科学研究所
	関西光科学研究所施記	安公開(木津地区)	
5	11-13-NOV-2021	オンライン開催(出展)	主催:けいはんな R&D フェア 2020 実行委員 会、関西文化学術研究都市推進機構
	京都 R&D フェア 202	21、けいはんなビジネスメッセ	2021
6	3-DEC-2021	オンライン開催	主催:関西文化学術研究都市推進機構
	けいはんなサイエンス	スカフェ(第 88 回)	
7	4-FEB-2022	オンライン開催	主催: RIKEN-RAP & QST- KPSI
	第5回RIKEN-RAP&	QST-KPSI ジョイントセミナー	

【播磨地区】

1	14-16-JUN-2021	つくば国際会議場+オンライ	主催:公益社団法人日本顕微鏡学会
		ン開催	
	公益社団法人日本顕後	微鏡学会 2021 年度第 77 回学術	講演会「S-8 ナノテクノロジープラットフォー
	ム・微細構造解析プ	ラットフォーム 10 年間の活動と	成果」シンポジウム
2	11-14-JUL-2021	SPring-8	主催:兵庫県立大学理学部/大学院理学研究
		(兵庫県佐用郡佐用町)	科、関西学院大学理学部・工学部・生命環境学
			部・大学院理工学研究科、東京大学 放射光分
			野融合国際卓越拠点、岡山大学、大阪大学未来

	1	1	
			戦略光科学連携センター・蛋白質研究所・核物 理研究センター、茨城大学大学院理工学研究 科、(公財)高輝度光科学研究センター(JASRI)、 理化学研究所 放射光科学研究センター、日本 原子力研究開発機構 物質科学研究センター、
			量子科学技術研究開発機構 関西光科学研究所
	第 21 回 SPring-8 夏の)学校	
3	16-SEP-2021	物質・材料研究機構手理地区	主催・国立研究開発注し物質・材料研究機構す
5	10 SEI 2021	10頁 初初初760歳時一先地区	
	文部科学省ナノテク /	ノロジーブフットフォーム令和:	3 年度利用成果発表会
4	17-SEP-2021	物質・材料研究機構千現地区	主催:国立研究開発法人物質・材料研究機構ナ
		+オンライン開催	ノテクノロジープラットフォームセンター
	会和3年度文部科学(省十ノテクノロジープラットフ-	オーム学生研修プログラム成里登表会
			ムーンナ工研修ションノン成本元表式
-	14 007 2021		
2	14-001-2021	オンフィン開催	王催: SPring-8 ユーザー協同体 (SPRUC)、(公
			財) 高輝度光科学研究センター、理化学研究所
			放射光科学研究センター
	合和3年度文部科学4	省ナノテクノロジープラットフィ	オーム事業 IAEA & OST 微細構造解析プラット
	フォーム放射光設備新	日田講羽合 一反広・合成レバル	
(
6	8-10-NOV-2021	幕張メッセ国際展示場(十葉	王傕:QST 微細構造解析ファットフォーム、
		県千葉市)	JAEA 微細構造解析プラットフォーム
	Japan Analytical & Sci	entific Instruments Show (JASIS)	2021
7	24-26-NOV-2021	九州大学医学部百年講堂	主催,一般社団法人日本分析機器工業会 一般
,			上语: 从正因因入口//··// // // // // // // // // // // //
	口中的他位当人一位		11回伝八日平村子城谷励云
	日本頭僦鏡子会 弟	64 回ンンホンワム 又部科学者	1) ノアクノロシーノフットノオーム (線) 一個
	析ブフットフォーム	ブース展示	
8	16-DEC-2021	オンライン開催	主催:日本顕微鏡学会
	文部科学省ナノテク	ノロジープラットフォーム JAI	EA&OST 微細構造解析プラットフォーム合同地
	城セミナー - 最新の	の雷子状能・表面反応・結晶構	告解析 _
0	$\frac{1}{3}$ $\frac{1}{2}$ $\frac{1}{3}$ $\frac{1}$	一 市 古 十 尚 木 卿 と 」 ン パ マ 」 ナ	
9	7-9-JAIN-2022	東京人子本郷キャンハス+オ	土催:JAEA 劔神博宣解ケノフットノオーム、
		ンフイン開催	QST 微細構造解析ブラットフォーム
	第35回日本放射光学	会年会・放射光科学合同シンポ	ジウム
10	26-28-JAN-2022	東京ビッグサイト+オンライ	主催・第35回日本放射光学会年会・放射光科
10	20 20 011 02022		一 二 に 、 、 、 、 こ こ 日 (「 八 川 川 川 一 一 八 川 川 一 一 八 川 川 一 一 一 八 川 川 一 一 一 八 川 川 一 一 一 八 川 川 一 一
			手口向シンホンリン組織安良去
	弗 20 回国際アノアク	ノロシー総合展・技術会議(na	no tech 2021)
		1	
11	28-JAN-2022	東京ビッグサイト+オンライ	主催: nano tech 2022 実行委員会
		ン開催	
		ノロジープラットフォーム第20	回ナノテカノロジー総合シンプポジウム
	JAPANNANO 2022	リノリクノロシーノリットノオ リーク ショー ニュロート ショー	$-\Delta M G \langle \mathcal{F} \mathcal{F} \mathcal{F} \mathcal{F} \mathcal{D} \mathbf{X} \rangle \mathcal{F} \mathcal{F} \mathcal{F} \mathcal{F} \mathcal{F} \mathcal{F} \mathcal{F} \mathcal{F}$
	~マアリアル先端リ	サーティンファの目指すもの~	
12	3-MAR-2022	東京大学本郷キャンパス+オ	主催:文部科学省ナノテクノロジープラット
		ンライン開催	フォーム、国立研究開発法人物質・材料研究
			機構ナノテクノロジープラットフォームセン
	微細構造解析プラッ	トフォームシンボジウム(10 年	の総括)
1			

13	4-MAR-2022	オンライン開催	主催:国立研究開発法人物質・材料研究機構
			微細構造解析プラットフォーム
	令和3年度文部科学行	省ナノテクノロジープラットフォ	オーム事業京大・JAEA・QST 微細構造解析プラ
	ットフォーム合同地均	或セミナー ―最新の微細構造	・状態解析—
14	8-MAR-2022	物質·材料研究機構千現地	主催:京大微細構造解析プラットフォーム最先
		区+オンライン開催	端構造観察·計測共用拠点、QST 微細構造解析
			プラットフォーム、JAEA 微細構造解析プラッ
			トフォーム
	文部科学省ナノテク /	ノロジープラットフォーム令和	3年度総会
15	9-MAR-2022	物質·材料研究機構千現地	主催:国立研究開発法人物質・材料研究機構ナ
		区+オンライン開催	ノテクノロジープラットフォームセンター
	令和3年度文部科学	省ナノテクノロジープラットファ	ォーム技術スタッフ交流プログラム報告会

KPSI セミナー(第79回~85回)

QST 木津地区では国内外の著名な研究者をお招きして学術的に最先端の 専門的なセミナーを開催しています。今年度は人数制限した会場から Web 中継も行うハイブリッドセミナーを中心に合計 7 回開催しました。開催に あたっては KPSI Web サイトやけいはんなメーリングリストを活用し、関 西研内外に開催案内を行っています。また報告については Web サイト、関 西研だより等を活用しています。(参加費無料・事前登録制)



高部英明教授

Web サイト: https://www.qst.go.jp/site/kansai-topics/29853.html

79	27-MAY-2021	Prof. Hideaki Takabe	National Taiwan University		
	Hawking radiation fro = Grasping physics in	Hawking radiation from true and analog black holes = Grasping physics intuitively and thinking about analog black hole experiments with ultra-intense lasers =			
80	28-SEP-2021	染川 智弘	(公財)レーザー技術総合研究所、大阪大 学レーザー科学研究所		
	海底開発における環	境影響評価に向けた水中レーザー	リモートセンシング技術の開発		
81	8-OCT-2021	Associate Prof. Jiro Itatani	ISSP, The Univ. of Tokyo		
	Intense ultrashort-pulse sources for materials science				
82	12-OCT-2021	Associate Professor Tomomasa Ohkubo	Tokyo University of Technology,		
	Development of Solar-Pumped Laser				
83	18-OCT-2021	Prof. Kenichi L. Ishikawa	The University of Tokyo		
	Building Science and Theory of Laser-Matter Interaction toward Super-Smart Society				
84	10-DEC-2021	佐々木 茂美	広島大学		
	量子ビームが運ぶ光	の軌道角運動量 光の軌道角運動量	を持つX線ビームで何が出来るか?		

85	20-JAN-2022	紀井 俊輝	京都大学
	バルク超伝導体の新	D新しい応用 遮蔽電流制御による高度な磁場分布デザイン	



左から、第80回 染川智弘博士、第81回 板谷治郎先生、第82回 大久保友雅先生



左から、第83回 石川顕一博士、第84回 佐々木茂美先生、第85回 紀井俊輝先生

QST 播磨セミナー(第 22 回~第 30 回)

播磨地区では QST 内の研究者や国内外の著名な研究者によるセミナーを開催しています。2021 年度は新型コロナウイルス感染症防止対策のため、オンライン開催としました。

22	21-APR-2021	Dr. Ayumi Shiro	High Pressure Science and Stress Research Group, QST	
	Tumor spheroid destru monochromatic X-rays	ction by gadolinium-loaded nanopartic s	cles irradiated with synchrotron radiation	
23	26-MAY-2021	Dr. Yoshinori Katayama	Kansai Photon Science Institute, QST	
	Overview of Advanced	Research Infrastructure for Materials	Science Project	
24	23-JUN-2021	Dr. Kenji Tsutsui	Condensed Matter Theory Group, QST	
	Antiphase Oscillations of the Time-Resolved Spin Structure Factor in Antiferromagnetic Mott Insulators			
25	14-JUL-2021	Dr. Mao Ye	State Key Laboratory of Functional Materials for Informatics, Shanghai Institute of Microsystem and Information Technology, Chinese Academy of Sciences	
	Recent Progress and Activities of the Soft-X-ray and Vacuum-Ultraviolet Beamlines at Shanghai Synchrotron Radiation Facility			

26	21-JUL-2021	Dr. Morihisa Saeki	Tokai Quantum Beam Science Center, Takasaki Advanced Radiation Research Institute, QST	
	Dispersive XAFS Study on Laser-induced Photoreduction of Rhodium(III) Complexes: Information Extraction of Rh Intermediate by MCR-ALS Analysis			
27	25-AUG-2021	Dr. Hideaki Iwasawa Dr. James Harries	Magnetism Research Group, QST Coherent X-ray Research Group, QST	
	Remote experiments at	t domestic and international synchrotro	on radiation facilities	
28	22-SEP-2021	Dr. Hiroyuki Saito	High Pressure Science and Stress Research Group, QST	
	 Hydrogen storage with earth abundant metals Current situation of data structuring in Materials Advanced Research Infrastructure project 			
29	27-OCT2021	Dr. Tetsuro Ueno	Magnetism Research Group, QST	
	Automated stopping cr	iterion for spectral measurement with	active learning	
30	30-NOV-2021	Dr. Takuji Nomura	Condensed Matter Theory Group, QST	
	Theory of L-shell x-ray magnetic circularly polarized emission: New optical sum rules			

関西光科学研究所2021年度年報 KPSI Annual Report 2021

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